PEGASIS: Practical Effective Class Group Action using 4-Dimensional Isogenies

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Abstract. In this paper, we present the first practical algorithm to compute an effective group action of the class group of any imaginary quadratic order \mathcal{O} on a set of supersingular elliptic curves primitively oriented by \mathcal{O} . Effective means that we can act with any element of the class group directly, and are not restricted to acting by products of ideals of small norm, as for instance in CSIDH. Such restricted effective group actions often hamper cryptographic constructions, *e.g.* in signature or MPC protocols.

Our algorithm is a refinement of the Clapoti approach by Page and Robert, and uses 4-dimensional isogenies. As such, it runs in polynomial time, does not require the computation of the structure of the class group, nor expensive lattice reductions, and our refinements allows it to be instantiated with the orientation given by the Frobenius endomorphism. This makes the algorithm practical even at security levels as high as CSIDH-4096. Our implementation in SageMath takes 1.5s to compute a group action at the CSIDH-512 security level, 21s at CSIDH-2048 level and around 2 minutes at the CSIDH-4096 level. This marks the first instantiation of an effective cryptographic group action at such high security levels. For comparison, the recent KLaPoTi approach requires around 200s at the CSIDH-512 level in SageMath and 2.5s in Rust.

1 Introduction

The simplicity and flexibility of the Diffie-Hellman key-agreement protocol [26] has made it ideally suited to create more advanced protocols, such as public-key encryption [28], digital signatures [60], password-authenticated key exchange [35], threshold encryption [25], threshold signatures [34, 33], updatable encryption [11] and many more. These protocols are all based on the computational efficiency and commutativity of group exponentiation, combined with the computational hardness of the discrete logarithm problem.

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The Clapoti Cult.

Due to Shor's algorithm [61], the discrete logarithm problem is not quantum hard. To construct a post-quantum analogue of the Diffie-Hellman protocol, we need to remove the underlying group structure. One way to achieve this is to use a commutative group action instead. The vectorisation and parallelisation problems, which are analogues of the discrete-logarithm and computational Diffie-Hellman problems respectively, are assumed to be (superpolynomially, but sub-exponentially [37]) hard, even for quantum computers. Furthemore, both problems are equivalently hard for quantum computers [31, 40, 32]. Using group actions, it is possible to construct post-quantum analogues of the advanced protocols given above: public-key encryption [41], digital signatures [8], passwordauthenticated key-exchange [1], threshold encryption and signatures [24] and updatable encryption [38].

A group action is said to be *effective* (EGA) if, amongst other conditions, the action can be computed in polynomial time [2]. That is, if G acts on X, we have a polynomial-time algorithm that computes gx for any g in G and x in X. If we only have a polynomial-time algorithm that can compute g_ix for $\{g_1, \ldots, g_n\}$, a set of (polylogarithmically many) generators of G, the action is said to be a *restricted effective* group action (REGA).

The most prominent instantiation of a REGA is given by the action of the class group of an imaginary quadratic order on a certain set of oriented elliptic curves, with the CSIDH protocol [14] (or its variant CSURF [13]) being the most efficient. In CSIDH, the orientation is given by the Frobenius endomorphism, and the set of oriented curves simply consists of the supersingular curves defined over \mathbb{F}_p , up to \mathbb{F}_p -isomorphism. Since we can only efficiently evaluate the action of ideals of smooth norm, CSIDH is indeed a REGA and not an EGA. This often results in difficulties in constructing post-quantum variants of protocols based on Diffie-Hellman: many of the previously mentioned examples really require an EGA, while other examples, such as the signature scheme SeaSign [22], resort to some form of rejection sampling to not leak information about the secret (as a workaround for being a REGA).

A three step approach to turn a REGA into an EGA was given by the signature scheme CSI-FiSh [8]: the first step computes the structure of the class group (which can be done in quantum polynomial time, but classically takes subexponential time), the second step computes a (very) short basis of the relation lattice (which takes exponential time, even quantumly) and the third step expresses any element as a product of the generators with small exponents, using the short basis. Step 1 has been done in practice for CSIDH-512, but further research [7, 10, 50] concludes that using a 512 bit prime for CSIDH does not achieve NIST level 1 security, with the latter two sources claiming that primes of more than 2000 bits are required. Since the computation of the class group is infeasible for such large primes, the primitive SCALLOP [21] and improvements [15, 3] avoided such computation by using a (large prime conductor) suborder of an order with small discriminant, so that the class group structure comes for free. This approach however does not fundamentally solve steps 2 and 3 (see https://yx7.cc/blah/2023-04-14.html for an informal, but detailed

discussion of this issue), and the highest level achieved, in [3] is equivalent to CSIDH-1536, for which a single group action takes almost 12 minutes using a C++ implementation. This approach is therefore completely hopeless to instantiate practical effective group actions at higher security levels (even using a quantum computer).

Our contributions. We describe a practical algorithm to compute the group action of any element in the class group of an imaginary quadratic order \mathcal{O} on a set of supersingular elliptic curves over $\overline{\mathbb{F}}_p$ primitively oriented by \mathcal{O} . Our algorithm works for all orders of discriminant $\Delta_{\mathcal{O}}$ all the way up to $|\Delta_{\mathcal{O}}| \approx p$ and in particular, applies in the CSIDH/CSURF setting, where the orientation is given by Frobenius, and we have $\Delta_{\mathcal{O}} = -p$ or $\Delta_{\mathcal{O}} = -4p$. This way, we obtain the first EGA at security levels equivalent to CSIDH-2048 and CSIDH-4096. We implement the algorithm in SageMath, and show that it achieves a highly practical runtime, which scales well to higher security levels. Not only does our algorithm allow us to easily instantiate security levels far beyond what previous EGA instantiations were able to; the timings of our proof-of-concept SageMath implementation also outperform all other EGA instantiations (even compared to their highly optimized C++/Rust implementations) at corresponding security levels. Find our implementation at https://github.com/pegasis4d.

Technical overview. Our work is based on the Clapoti framework [47], and proceeds by embedding the isogeny we wish to evaluate in an isogeny in dimension 4. In order to evaluate the action given by the ideal $\mathfrak{a} \subseteq \mathcal{O}$ on an elliptic curve E using only one 4-dimensional 2^e -isogeny, the Clapoti framework requires finding two other integral ideals $\mathfrak{b}, \mathfrak{c}$ equivalent to \mathfrak{a} such that

$$uN(\mathfrak{b}) + vN(\mathfrak{c}) = 2^e,\tag{1}$$

with the restriction that u, v are integers that can be written as the sum of two squares (in particular, they must be easy to factorise), and where $2^e < p$ (or more accurately, the full torsion group $E[2^e]$ should be defined over $\mathbb{F}_{p^{2k}}$ for a small k).

Several issues arise when trying to solve the above norm equation for $|\Delta_{\mathcal{O}}| \approx p$: even without the restriction on u and v being sums of squares, sometimes the equation simply has no solution. Indeed, since the smallest ideals equivalent to \mathfrak{a} are already expected to have norm around \sqrt{p} , and a solution is guaranteed to exist only when $N(\mathfrak{b})N(\mathfrak{c}) < 2^e$, this contradicts the requirement $2^e < p$. As such, imposing the extra condition for both u and v to be sums of squares further lowers the probability of finding a solution.

The main insight is that we can derive a related equation which is much easier to solve by exploiting the fact we can always efficiently compute small degree isogenies. We use this insight in two ways: first, we write $\mathbf{b} = \mathbf{b}_k \mathbf{b}_e$ and $\mathbf{c} = \mathbf{c}_k \mathbf{c}_e$ where \mathbf{b}_e and \mathbf{c}_e are ideals of smooth norm and compute the isogenies $\varphi_{\mathbf{b}_e} : E \to E_1$ and $\varphi_{\mathbf{c}_e} : E \to E_2$, simply using Elkies' algorithm [29] or Vélu's algorithm [62]. We can then apply the Clapoti framework to $E_1 \times E_2$ instead of $E \times E$, resulting in the equation

$$uN(\mathfrak{b}_k) + vN(\mathfrak{c}_k) = 2^e,$$

where now the product $N(\mathfrak{b}_k)N(\mathfrak{c}_k)$ is typically much smaller than 2^e since we have removed the norms of the ideals \mathfrak{b}_e and \mathfrak{c}_e .

Second, we use the same insight to relax the conditions on u and v: instead of requiring that each can be written as a sum of squares, we again allow to take out smooth factors g_u and g_v such that $u = g_u(x_u^2 + y_u^2)$ and $v = g_v(x_v^2 + y_v^2)$. Note that integers containing prime factors (to an odd power) which are 3 mod 4 can never be written as a sum of squares, so allowing factors like 3, 7, 11, 19, ... to be taken out of u and v drastically increases the probability that both u/g_u and v/g_v are sums of squares. Applying the above, allows us to solve the (relaxed) norm equation efficiently, even at the highest security levels.

With u and v of this form, we are able to efficiently construct a 2-dimensional u-isogeny $\Phi_u : E_1^2 \to E_u^2$ and a 2-dimensional v-isogeny $\Phi_v : E_2^2 \to E_v^2$. We then compute a 4-dimensional 2^e -isogeny $F : E_u^2 \times E_v^2 \to E_a^2 \times E'^2$ that "embeds" Φ_u , $\Phi_v, \varphi_{\mathfrak{b}_k} : E_1 \to E_{\mathfrak{a}}$ and $\varphi_{\mathfrak{c}_k} : E_2 \to E_{\mathfrak{a}}$, where $E_{\mathfrak{a}} := \mathfrak{a} \cdot E$. Using level 2 theta coordinates as in [18], the computation of F is relatively fast. We finally extract the resulting curve $E_{\mathfrak{a}}$ from the codomain of F. For very high security levels, we remark that it is helpful to relax the conditions on u, v and allow them to be arbitrary positive integers, at the cost of using more dimension 4 isogenies (see Sections 4.2 and 6.4).

Related work. A recent paper by Panny, Petit and Stopar [49] has also instantiated a primitive KLaPoTi based on Clapoti. KLaPoTi differs from PEGASIS in several aspects. The main ones are the way Equation (1) is solved, the dimension of the isogeny used in the evaluation of the group action and the size of the base prime p. KLaPoTi relies on the KLPT algorithm [36] to solve Equation (1), which now requires u, v to be perfect squares. The fact that u and v are perfect squares allows to perform the group action computation using dimension 2 isogenies only. Nevertheless, the relatively large size of the solution obtained by this KLPT approach requires $|\Delta|^3 < 2^e$ where Δ is the discriminant of the order \mathcal{O} . This implies that KLaPoTi uses a base prime p which is 3 times larger than the one used in CSIDH and PEGASIS, and uses an order \mathcal{O} of discriminant Δ with $|\Delta|^3 < 2^e < p$. This highly affects the performance and the key sizes in KLaPoTi. Even though PEGASIS performs the group action using 4-dimensional isogenies, the fact that $|\Delta| \approx 2^e \approx p$ in PEGASIS enables a group action computation which is more efficient. For example, to achieve similar security to CSIDH-512, KLaPoTi uses a prime p of 1536 bits and a single group action computation takes approximately 3 minutes [49, Section 7.1] in SageMath, while PEGASIS uses a prime of about 512 bits and a single group action computation takes approximately 1.5 seconds.

Organisation of paper. The rest of the paper is organized as follows. In Section 2 we recall the necessary background material. In Section 3 we describe our whole algorithm for general orientations. In Section 4, we detail an important subroutine, namely how to compute an isogeny of prescribed degree in dimension 2, before we in Section 5 specialise our algorithm for the CSIDH orientation. Finally, in Section 6 we detail some implementation choices before we present our timings.

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2 Preliminaries

For a general introduction to isogenies, especially in the context of cryptography, we refer the reader to [30].

2.1 Isogeny class-group action on oriented curves

Classic results [63, Th. 4.5] tell us that the class-group $\operatorname{Cl}(\mathcal{O})$ of an imaginary quadratic order \mathcal{O} acts freely and transitively on the set $\operatorname{Ord}_q(\mathcal{O})$ of ordinary elliptic curves E/\mathbb{F}_q with endomorphism ring $\operatorname{End}(E) \cong \mathcal{O}$. In [16], Colò and Kohel introduced a similar action, in which $\operatorname{Cl}(\mathcal{O})$ acts on a set of supersingular elliptic curves E which contain \mathcal{O} as a subring of $\operatorname{End}(E)$. We briefly recount this construction, together with results from [46].

Let $\mathcal{O} = \mathbb{Z}[\alpha]$ be an imaginary quadratic order and E/\mathbb{F}_{p^n} an elliptic curve. We call an injective morphism of rings $\iota: \mathcal{O} \hookrightarrow \operatorname{End}(E)$ an \mathcal{O} -orientation and say it is primitive if $\iota(\alpha) = \omega$ is a primitive element of $\operatorname{End}(E)$ viewed as a lattice, *i.e.* ω does not factor through a (non-trivial) scalar multiplication. We call the pair (E, ι) a (primitively) \mathcal{O} -oriented elliptic curve.

Let (E, ι) be primitively \mathcal{O} -oriented and \mathfrak{a} an integral invertible ideal of \mathcal{O} with norm prime to p. Let $\varphi_{\mathfrak{a}} \colon E \to E_{\mathfrak{a}}$ be the isogeny with kernel

$$E[\mathfrak{a}] = \bigcap_{\sigma \in \mathfrak{a}} \ker(\iota(\sigma))$$

and degree N(\mathfrak{a}). Now consider the map $\mathcal{O} \to \operatorname{End}(E_{\mathfrak{a}}); \gamma \mapsto \varphi_{\mathfrak{a}}\iota(\gamma)\widehat{\varphi_{\mathfrak{a}}}$. It is an injective morphism of additive groups, but sends $1 \mapsto [\operatorname{deg}(\varphi_{\mathfrak{a}})]$ and so is not a morphism of rings. We fix this by first noting that $\varphi_{\mathfrak{a}}\iota(\gamma)\widehat{\varphi_{\mathfrak{a}}}$ always factors

through $[\deg(\varphi_{\mathfrak{a}})]$ and then defining $(\varphi_{\mathfrak{a}})_*(\iota) := \varphi_{\mathfrak{a}}\iota(\gamma)\widehat{\varphi_{\mathfrak{a}}}/[\deg(\varphi_{\mathfrak{a}})]$ to obtain a morphism of rings and thus a well-defined \mathcal{O} -orientation. Indeed, by diagramchasing one verifies that $\ker(\varphi_{\mathfrak{a}})$ is invariant under $\omega = \iota(\alpha)$, demonstrating that $\varphi_{\mathfrak{a}}\iota(\gamma)\widehat{\varphi_{\mathfrak{a}}}(E[\deg(\varphi_{\mathfrak{a}})]) = \{0\}$ for all γ in $\mathbb{Z}[\alpha]$, and that $\varphi_{\mathfrak{a}}\iota(\gamma)\widehat{\varphi_{\mathfrak{a}}}$ factors through $[\deg(\varphi_{\mathfrak{a}})]$. In fact, $(\varphi_{\mathfrak{a}})_*(\iota)$ is a *primitive* orientation [46, Prop. 3.5].

We say that the isogenies $\varphi_{\mathfrak{a}} \colon (E, \iota) \to (E', (\varphi_{\mathfrak{a}})_*(\iota))$ are \mathcal{O} -oriented. Likewise, an isomorphism $\varphi \colon E \to E'$ is called an \mathcal{O} -isomorphism $(E, \iota) \to (E', \iota')$ if $\iota' = \varphi_*(\iota) \coloneqq (\gamma \mapsto \varphi\iota(\gamma)\hat{\varphi})$. Using this terminology, we define $\mathrm{SS}_{p^n}^{\mathrm{pr}}(\mathcal{O})$ to be the \mathcal{O} -isomorphism classes of primitively \mathcal{O} -oriented supersingular elliptic curves defined over \mathbb{F}_{p^n} . This set is non-empty if and only if p is not split in $K = \mathcal{O} \otimes_{\mathbb{Z}} \mathbb{Q}$ and does not divide the conductor of \mathcal{O} in \mathcal{O}_K [46, Prop. 3.2].

We note that every class $[\mathfrak{a}]$ in the class-group $\operatorname{Cl}(\mathcal{O})$ is represented by an invertible integral ideal \mathfrak{b} with norm $\operatorname{N}(\mathfrak{b})$ coprime to p [17, Cor. 7.17]. Moreover, it is clear that principal ideals $\gamma \mathcal{O}$ correspond to endomorphisms $\varphi_{\gamma \mathcal{O}} = \iota(\gamma)$. Now the map

$$\operatorname{Cl}(\mathcal{O}) \times \operatorname{SS}_{p}^{\operatorname{pr}}(\mathcal{O}) \to \operatorname{SS}_{p}^{\operatorname{pr}}(\mathcal{O}) \qquad [\mathfrak{a}], (E,\iota) \mapsto \mathfrak{a} \cdot (E,\iota) := (E_{\mathfrak{a}}, (\varphi_{\mathfrak{a}})_{*}(\iota))$$

is well-defined; and when when $SS_p^{pr}(\mathcal{O})$ is non-empty, the action is free with at most two orbits [46, Prop. 3.3, Th. 3.4]. Restricting to one of these orbits, we obtain a free and transitive group action.

The vectorisation problem of this action - given $(E, \iota), (E', \iota')$ compute $[\mathfrak{a}]$ such that $\mathfrak{a} \cdot (E, \iota) = (E', \iota')$ - is thought to be quantum resistant. This property led to several cryptographic constructions based on ideal class group actions [14, 16, 21, 15] which define (restricted) effective group actions, as mentioned in Section 1. The first one to be introduced was CSIDH (Commutative Supersingular Isogeny Diffie-Hellman) [14] where $\mathcal{O} = \mathbb{Z}[\sqrt{-p}]$ and the orientation induces an isomorphism $\mathcal{O} \cong \operatorname{End}_{\mathbb{F}_p}(\mathcal{O})$, mapping $\sqrt{-p}$ to Frobenius endomorphism of $(x, y) \mapsto (x^p, y^p)$ for all $E \in \operatorname{SSp}^{\operatorname{pr}}(\mathcal{O})$. Even though these schemes are thought to be quantum resistant, the best known quantum attack against the underlying vectorisation problem is a subexponential algorithm due to Kuperberg [37]. For that reason, there is an uncertainty on the parameters required to ensure a sufficient security level for CSIDH (and other class group action schemes). As mentioned in Section 1, conservative sources [7, 10, 50] estimate that a 2000 bit prime p is required to ensure a NIST-1 security level for CSIDH.

2.2 Polarised isogenies between abelian varieties

Every Abelian variety A has a dual \widehat{A} of the same dimension and every isogeny $\varphi: A \to B$ has a dual $\widehat{\varphi}: \widehat{B} \to \widehat{A}$ of the same degree. If there exists a special kind of isogeny $\lambda_A: A \to \widehat{A}$ called a *polarisation*, we say (A, λ_A) is a *polarised* Abelian variety (PAV). When λ_A is an isomorphism, we say it is a *principal* polarisation and call A a *principally* polarised Abelian variety (PPAV). Elliptic curves are exactly the 1-dimensional PPAVs; they are canonically principally polarised.

Definition 1 (Polarised Isogenies). Let $\varphi: (A, \lambda_A) \to (B, \lambda_B)$ be an isogeny between PAVs. If there exists an integer d such that $\widehat{\varphi}\lambda_B\varphi = [d]\lambda_A$, we say

that φ is (λ_A, λ_B) -polarised with polarised degree $\deg_p(\varphi) = d$. When A, B are PPAVs we define the polarised dual of φ to be $\tilde{\varphi} = \lambda_A^{-1} \hat{\varphi} \lambda_B \colon (B, \lambda_B) \to (A, \lambda_A)$. Then φ is (λ_A, λ_B) -polarised with degree d if $\tilde{\varphi} \varphi = [d]$ in End(A).

 (λ_A, λ_B) -polarised isogenies of polarised degree d are characterised by an isotropy condition of their kernel.

Lemma 2. Let $\varphi: (A, \lambda_A) \to (B, \lambda_B)$ be a (λ_A, λ_B) -polarised isogeny with polarised degree d such that $\operatorname{char}(k) \nmid d$ and let $g := \dim(A)$. Then $\operatorname{ker}(\varphi)$ is a subgroup of cardinality d^g of A[d], isotropic with respect to the Weil pairing $e^{[d]\lambda_A}$ associated to the polarisation $[d]\lambda_A: A \to \widehat{A}$ (as defined in [39, p. 135]).

Conversely, if A is a PPAV of dimension g and $\varphi: A \to B$ has kernel $\ker(\varphi) \subseteq A[d]$ of cardinality d^g and isotropic for $e^{[d]\lambda_A}$. Then there exists a unique principal polarisation λ_B on B such that $\varphi: A \to B$ is a (λ_A, λ_B) -polarised isogeny with polarised degree d.

Proof. Suppose that φ is a (λ_A, λ_B) -polarised isogeny with polarised degree d. Let $x, y \in \ker(\varphi)$ and $y' \in B$ such that $y = \tilde{\varphi}(y')$ (which exists by surjectivity of $\tilde{\varphi}$ and belongs to B[d] since $\varphi \circ \tilde{\varphi} = [d]$). Then, by the definition of $e^{[d]\lambda_A}$ and the usual properties of the Weil pairing [39, Lemma 16.2],

$$e^{[d]\lambda_A}(x,y) = e_d(x,\lambda_A(y)) = e_d(x,\lambda_A \circ \widetilde{\varphi}(y')) = e_d(x,\widehat{\varphi} \circ \lambda_B(y'))$$
$$= e_d(\varphi(x),\lambda_B(y')) = 1,$$

since $\varphi(x) = 0$. This proves that $\ker(\varphi)$ is isotropic.

Conversely, suppose that $\ker(\varphi) \subseteq A[d]$ of cardinality d^g and isotropic for $e^{[d]\lambda_A}$. Then [39, Proposition 16.8] ensures the existence of a polarisation λ_B on B such that $\widehat{\varphi} \circ \lambda_B \circ \varphi = [d]\lambda_A$. Since $\deg(\widehat{\varphi}) = \deg(\varphi) = |\ker(\varphi)| = d^g$ and $\deg([d]) = d^{2g}$, it follows that $\deg(\lambda_B) = \deg(\lambda_A) = 1$ by multiplicativity of the degrees so λ_B is a principal polarisation. This completes the proof. \Box

For brevity, we will drop mention of the polarisations when they are clear from context, and say *d-isogeny* to mean an isogeny with polarised degree *d*. In particular, it is implicit that the corresponding polarisation to a PAV denoted *A* is λ_A . Between elliptic curves, the principal polarisation being canonical, the dual $\hat{\varphi}$ and polarised dual $\tilde{\varphi}$ of an isogeny can be identified and will both be denoted by $\hat{\varphi}$; and the notions of degree and polarised degree are also the same.

We recall a standard result about factoring isogenies between PPAVs, which we will need to recall the Clapoti construction in the more general context of [48, Remark 2.10].

Lemma 3 (Factoring Isogenies between PPAVs). Let A, B be PPAVs defined over a field k and $\varphi: A \to B$ be a d-isogeny between them. For every pair of positive coprime integers d_1, d_2 each being coprime to char(k) satisfying $d_1d_2 =$ d, there exists a PPAV C, a d_1 -isogeny $\varphi_1: A \to C$ and a d_2 -isogeny $\varphi_2: C \to B$, each uniquely defined up to isomorphism, such that $\varphi = \varphi_2 \circ \varphi_1$. *Proof.* Since d_1 and d_2 are coprime, we have $\ker(\varphi) = \ker(\varphi)[d_1] \oplus \ker(\varphi)[d_2]$ with $|\ker(\varphi)[d_i]| = d_i^g$ for $i \in \{1, 2\}$. Consider the isogeny $\varphi_1 \colon A \to C$ of kernel $\ker(\varphi)[d_1]$ and $\varphi_2 \colon C \to B'$ the isogeny of kernel $\varphi_1(\ker(\varphi))$. Then by construction $\ker(\varphi_2 \circ \varphi_1) = \ker(\varphi)$ so $\varphi_2 \circ \varphi_1 = \varphi$ up to postcomposition with an isomorphism $B \to B'$ by [43, Theorem 4, p. 74].

Since φ is a *d*-isogeny, Lemma 2 ensures that ker $(\varphi) \subseteq A[d]$ is isotropic for $e^{[d]\lambda_A}$ so ker $(\varphi_1) \subseteq A[d_1]$ is isotropic for $e^{[d_1]\lambda_A}$ by the usual properties of the Weil pairing [39, Lemma 16.1] and of cardinality d_1^g , as we saw. Using Lemma 2 again, we obtain a principal polarisation λ_C on C such that φ_1 is a (λ_A, λ_C) -polarised isogeny of polarised degree d_1 .

To conclude, we verify that φ_2 is a (λ_C, λ_B) -polarised isogeny of polarised degree d_2 . Substituting into $\widehat{\varphi}\lambda_B\varphi = [d_1d_2]\lambda_A$, we obtain $\widehat{\varphi}_1\widehat{\varphi}_2\lambda_B\varphi_2\varphi_1 = [d_2d_1]\lambda_A = [d_2]\widehat{\varphi}_1\lambda_C\varphi_1$ and conclude that $\widehat{\varphi}_2\lambda_B\varphi_2 = [d_2]\lambda_C$. Indeed, isogenies are surjective so fh = gh implies f = g and by forming the dual (twice) we see that hf = hgalso implies f = g. Note that this last argument also proves the uniqueness of φ_1 and φ_2 . This completes the proof.

2.3 Kani's lemma and embedding isogenies into higher degree

If $(\varphi_{ji})_{ij}$ is a matrix of polarised isogenies $\varphi_{ij} \colon A_i \to B_j$ between PPAVs, then its polarised dual is given by $(\widetilde{\varphi_{ij}})_{ij}$. Whether or not a 2 × 2 matrix of polarised isogenies is a polarised isogeny is the content of Kani's lemma.

Lemma 4 (Kani, [53, Lemma 2.1]). Let A_1, A_2, B_1, B_2 be PPAVs defined over k and let $\varphi_{ij}: A_i \to B_j$ be polarised d_{ij} -isogenies. The map

$$\Phi = \begin{pmatrix} \varphi_{11} & \varphi_{21} \\ \varphi_{12} & \varphi_{22} \end{pmatrix} : A_1 \times A_2 \to B_1 \times B_2$$

is a polarised isogeny if and only if $d_{11} = d_{22}, d_{12} = d_{21}$ and the corresponding square S_{Φ}

$$\begin{array}{ccc} A_1 & \xrightarrow{\varphi_{11}} & B_1 \\ -\varphi_{12} & & & \downarrow \widetilde{\varphi_{21}} \\ B_2 & \xrightarrow{\varphi_{22}} & A_2 \end{array}$$

commutes. In such cases, $\deg_p(\Phi) = d_{11} + d_{12}$ and we call Φ the Kani-map corresponding to the Kani-square S_{Φ} . Further, if $d_{11} = d_{22}$ is prime to $d_{12} = d_{21}$, we call the Kani-square a (d_{11}, d_{12}) -isogeny diamond. Finally, when d_{11} is prime to d_{12} and $d = d_{11} + d_{12}$ is prime to the characteristic of k, then

$$\ker(\Phi) = \left\{ \left([d_{11}]x, \widetilde{\varphi_{21}}\varphi_{11}(x) \right) \mid x \in A[d] \right\}.$$

If Φ can be efficiently evaluated, then so can φ_{ij} , and Φ is said to *embed* φ_{ij} . In other words, Φ is an *efficient representation* of the φ_{ij} in the sense of the following definition. **Definition 5.** Let \mathscr{A} be an algorithm. Given a *d*-isogeny $\varphi : A \to B$ between PPAV of dimension *g* defined over \mathbb{F}_q , an *efficient representation* of φ with respect to \mathscr{A} is some data $D_{\varphi} \in \{0,1\}^*$ of polynomial size in *g*, $\log(d)$ and $\log(q)$ such that for all $P \in A(\mathbb{F}_{q^k})$, \mathscr{A} with input (D_{φ}, P) returns $\varphi(P)$ in polynomial time in *g*, $\log(d)$, *k* and $\log(q)$.

3 Our algorithmic approach

We now describe our algorithm for computing the class group action by a general orientation.

3.1 Applying Kani's lemma in dimension 4

The factorisation lemma (Lemma 3) gives us a strategy to embed arbitrary isogenies into higher dimension [48, Remark 2.10]. If $\varphi: A \to B, \psi: A \to C$ are isogenies between PPAVs with coprime polarised degrees, we can form a Kani-diamond by factoring the composition $\psi \tilde{\varphi}$ as $\tilde{\varphi'} \psi'$ whereby $\psi': B \to D$ and $\varphi': C \to D$ have polarised degrees $\deg_p(\varphi') = \deg_p(\varphi), \deg_p(\psi) = \deg_p(\psi')$. The kernel of the resulting Kani-map can be computed so long as $\psi \tilde{\varphi}$ is known on $B[\deg_p(\varphi) + \deg_p(\psi)]$.

When computing the class-group action of $[\mathfrak{a}]$ on a primitively oriented elliptic curve (E, ι) , the isogenies φ, ψ are taken to be $\widehat{\varphi}_{\mathfrak{b}}, \widehat{\varphi}_{\mathfrak{c}} : E_{\mathfrak{a}} \to E$ where $\mathfrak{b}, \mathfrak{c}$ are representatives of $[\mathfrak{a}]$. The composition $\widehat{\varphi}_{\mathfrak{c}}\varphi_{\mathfrak{b}} = \varphi_{\overline{\mathfrak{c}}\mathfrak{b}}$ is an endomorphism on E dictated by ι and so the kernel of the resulting 2-dimensional Kani-map can computed so long as the N(\mathfrak{b})+N(\mathfrak{c}) torsion is accessible. For practically efficient computation (using [20] for instance), we must find ideals $\mathfrak{b}, \mathfrak{c}$ representing $[\mathfrak{a}]$ such that N(\mathfrak{a}) + N(\mathfrak{b}) = 2^e , where $2^{e+2} \mid p+1$ in order to work over \mathbb{F}_{p^2} (the reason we need 2^{e+2} to divide p+1 is due to technicalities when using level 2 theta coordinates to compute higher dimensional isogenies). However, finding ideals $\mathfrak{b}, \mathfrak{c}$ satisfying this norm equation is not easy in practice and can only be done with standard techniques when p is much bigger than $|\operatorname{disc}(\mathcal{O})|$ (which is not suitable for CSIDH/CSURF, where $|\operatorname{disc}(\mathcal{O})| = p$ or 4p).

Goal: solve the norm equation:

$$u\mathbf{N}(\mathbf{b}) + v\mathbf{N}(\mathbf{b}) = 2^e \tag{2}$$

with $2^{e+2} | p+1$, $[\mathfrak{a}] = [\mathfrak{b}] = [\mathfrak{c}]$, u, v > 0, $gcd(uN(\mathfrak{b}), vN(\mathfrak{c})) = 1$, and such that we can build efficiently computable dimension d isogenies $\Phi_u : E^d \to A_u, \Phi_v : E^d \to A_v$ of polarised degrees u, v.

Indeed, the factoring lemma (Lemma 3) ensures that we have a $(uN(\mathfrak{b}), vN(\mathfrak{c}))$ -isogeny diamond



where $\Phi_{\mathfrak{b}} := \operatorname{diag}(\varphi_{\mathfrak{b}}), \Phi_{\mathfrak{c}} := \operatorname{diag}(\varphi_{\mathfrak{c}}) \colon E^d \to E^d_{\mathfrak{a}}, \Phi$ is an $uN(\mathfrak{b})$ -isogeny and Ψ is a $vN(\mathfrak{c})$ -isogeny. Hence, by Kani's lemma Lemma 4, we can embed the compositions $\Phi_{\mathfrak{b}} \circ \widetilde{\Phi}_u$ and $\Phi_{\mathfrak{c}} \circ \widetilde{\Phi}_v$ into a 2*d*-dimensional 2^e -isogeny $F : A_u \times A_v \to E^d_{\mathfrak{a}} \times A$.

Except when u and v are perfect squares as in [49] (which does not make the norm equation much easier to solve), we cannot expect Φ_u and Φ_v to be one dimensional. Using higher dimensions $d \geq 2$ appears necessary because it is difficult to construct (efficient representations of) outgoing 1-dimensional isogenies Φ_u, Φ_v of large given degree u, v on E without knowing the endomorphism ring $\operatorname{End}(E)$: the only tools we have at our disposal to produce arbitrary-degree isogenies all require higher dimensions. In fact, a variant of this problem (construct a large *prime* degree isogeny on E) is conjectured to be cryptographically hard [4, Problem 2].

In any case, Zarhin's trick would allow us to efficiently construct endomorphisms Φ_u, Φ_v of E^4 with the prescribed polarised degrees u, v. However, we consider computing the corresponding 8-dimensional isogeny F to be too expensive. Instead, we will see in Section 4 how to find solutions when d = 2. Consequently, we have to compute a 4-dimensional isogeny F, which can be done efficiently with level 2 theta coordinates [18]. As mentioned earlier, the algorithms to compute the 2^e -isogeny F require 2^{e+2} -torsion points, so we need $2^{e+2}|p+1$ instead of $2^e|p+1$ to be able to work over \mathbb{F}_{p^2} .

Unfortunately, there do not always exist positive integer solutions u, v to $uN(\mathfrak{b}) + vN(\mathfrak{c}) = 2^e$. Indeed, the *Frobenius coin problem* tells us that we are only guaranteed solutions when $2^e \geq N(\mathfrak{b})N(\mathfrak{c}) - N(\mathfrak{b}) - N(\mathfrak{c})$; on the other hand, when $\mathcal{O} = \mathbb{Z}[\sqrt{-p}]$ (as in CSIDH), the smallest representative of any class $[\mathfrak{a}]$ in $Cl(\mathcal{O})$ is expected to be of size roughly \sqrt{p} . So, recalling that $2^{e+2} | p+1$, we see that we rarely have guaranteed solutions u, v. The solution we propose to overcome this difficulty is to give some additional freedom on the choice of ideals $\mathfrak{b}, \mathfrak{c}$ by factoring out their smooth part.

3.2 Factoring out easy steps first

We will now describe how to compute the action of $[\mathfrak{a}]$ on (E, ι) in four dimensions. Throughout this section, $\mathfrak{b}, \mathfrak{c}$ are primitive representatives of $[\mathfrak{a}]$.

The idea Suppose we can factor $\mathfrak{b} = \mathfrak{b}_e \mathfrak{b}_k, \mathfrak{c} = \mathfrak{c}_e \mathfrak{c}_k$, so that the isogenies $\varphi_{\mathfrak{b}_e} : E \to E_1, \varphi_{\mathfrak{c}_e} : E \to E_2$ are efficiently computable (*e.g.* smooth degree) and so that we can find efficiently computable isogenies $\Phi_u : E_1^2 \to A_u, \Phi_v : E_2^2 \to A_v$ (*e.g.* integer-matrix endomorphisms) with polarised degrees u, v satisfying

$$u\mathbf{N}(\mathbf{b}_k) + v\mathbf{N}(\mathbf{c}_k) = 2^e.$$
(3)

Then we can consider the following $(uN(\mathfrak{b}), vN(\mathfrak{c}))$ -isogeny diamond



where Φ_I is the 2-dimensional isogeny diag (φ_I, φ_I) given by the action of $I \subseteq \mathcal{O}$. Then, Kani's lemma yields a 4-dimensional isogeny

$$F := \begin{pmatrix} \Phi_{\mathfrak{b}_k} \circ \widetilde{\Phi_u} \ \Phi_{\mathfrak{c}_k} \circ \widetilde{\Phi_v} \\ -\Psi & \widetilde{\Phi} \end{pmatrix} : A_u \times A_v \to E^2_{\mathfrak{a}} \times A, \tag{4}$$

with kernel

$$\ker(F) = \left\{ ([u\mathbf{N}(\mathfrak{b}_{k})]x, \Phi_{v}\widetilde{\Phi_{\mathfrak{c}_{k}}}\Phi_{\mathfrak{b}_{k}}\widetilde{\Phi_{u}}(x)) \mid x \in A_{u}[2^{e}] \right\}$$
$$= \left\{ ([u\mathbf{N}(\mathfrak{b}_{k})]x, \Phi_{v}\Phi_{\overline{\mathfrak{c}_{k}}\mathfrak{b}_{k}}\widetilde{\Phi_{u}}(x)) \mid x \in A_{u}[2^{e}] \right\}$$
$$= \left\{ ([\mathbf{N}(\mathfrak{b}_{k})]\Phi_{u}(y), \Phi_{v}\Phi_{\overline{\mathfrak{c}_{k}}\mathfrak{b}_{k}}(y)) \mid y \in E_{1}^{2}[2^{e}] \right\},$$
(5)

Since $\mathfrak{b}, \mathfrak{c}$ both represent $[\mathfrak{a}], \overline{\mathfrak{c}}\mathfrak{b} = \sigma \mathcal{O}$ is principal and we can evaluate $\varphi_{\overline{\mathfrak{c}}\mathfrak{b}} = \iota(\sigma)$. Having computed $\varphi_{\mathfrak{c}_e}$ and $\varphi_{\mathfrak{b}_e}$, we can evaluate

$$\mathbf{N}(\mathfrak{b}_e)\mathbf{N}(\mathfrak{c}_e)\varphi_{\overline{\mathfrak{c}_k}\mathfrak{b}_k} = \varphi_{\mathfrak{c}_e}\iota(\sigma)\widehat{\varphi_{\mathfrak{b}_e}}.$$
(6)

and compute ker(F) (more exactly points of 2^{e+2} -torsion generating ker(F) as required in [18]). We refer to Section 5.2 for more details on the kernel computation in the case of CSIDH.

The algorithm By construction, $uN(\mathfrak{b}_k) + vN(\mathfrak{c}_k) = 2^e$ has at least as many positive integer solutions u, v as $uN(\mathfrak{b}) + vN(\mathfrak{c}) = 2^e$. Indeed, every ideal appearing as \mathfrak{b}_k can also be chosen as \mathfrak{b} (with $\mathfrak{b}_e = 1$). More relevant, though, is that the norms of the ideals \mathfrak{b}_k are smaller, and so we hope for more u, v solutions.

The ideals $\mathfrak{b}_e, \mathfrak{c}_e$ should be chosen so that the corresponding isogenies $\varphi_{\mathfrak{b}_e}, \varphi_{\mathfrak{c}_e}$ are easy to compute, *e.g.* of smooth degrees. More precisely, letting \mathfrak{B} be a set

of small primes, we permit $\mathfrak{b}_e, \mathfrak{c}_e$ to be a product of ideals lying above the primes in \mathfrak{B} . Note that since $\mathfrak{b}_e, \mathfrak{c}_e$ are primitive, these primes will all be split in \mathcal{O} .

With this requirement we now proceed as follows (we refer to Section 5.1 for details).

- 1. Obtain a set of S of short representatives of $[\mathfrak{a}]$ (e.g. by Lagrange-reduction).
- 2. Factor each \mathfrak{b} in S as $\mathfrak{b} = \mathfrak{b}_e \mathfrak{b}_k$ maximising the norm of \mathfrak{b}_e .
- 3. Select a pair of ideals $\mathfrak{b} = \mathfrak{b}_e \mathfrak{b}_k$ and $\mathfrak{c} = \mathfrak{c}_e \mathfrak{c}_k$ from S with $N(\mathfrak{b}_k), N(\mathfrak{c}_k)$ coprime, prioritizing those with the smallest $N(\mathfrak{b}_k), N(\mathfrak{c}_k)$.
- 4. Enumerate the positive integer solutions u, v of $uN(\mathfrak{b}_k) + vN(\mathfrak{c}_k) = 2^e$.
- 5. Return a solution u, v if it allows for the construction of Φ_u, Φ_v ; else choose a different $\mathfrak{b}, \mathfrak{c}$ pair.

We briefly remark that this technique of splitting the isogeny computation into an easy 1-dimensional, followed by potentially expensive $(d \ge 2)$ dimensional computation also allows us to compute more isogenies Φ_u of prescribed polarised degree u. Indeed, the naïve approach of constructing Φ_u as a 2×2 integer-matrix

$$\begin{pmatrix} x_u - y_u \\ y_u & x_u \end{pmatrix} \tag{7}$$

only permits $u = x_u^2 + y_u^2$ to be a sum of two squares. Composing Φ_u as the diagonal of some one-dimensional isogeny of sufficiently smooth degree g_u (e.g. \mathfrak{B} -smooth), and then a matrix as above allows for polarised degrees $u = g_u(x_u^2 + y_u^2)$ (see Section 4.1).

3.3 Finding the codomain orientation

Once we have computed the 4-dimensional isogeny $F: A_u \times A_v \to E_a^2 \times A$ and extracted E_a from its codomain (following the approach presented in Appendix B in the case of CSIDH), we still have to compute the orientation $\iota_a := (\varphi_a)_*(\iota)$ on E_a . In the case of CSIDH, this orientation is naturally given by the Frobenius endomorphism and does not need to be computed. However, this orientation is not that easy to determine in general.

Given a generator $\alpha \in \mathcal{O}$ of \mathcal{O} ($\mathcal{O} = \mathbb{Z}[\alpha]$), we only have to compute an efficient representation of $\iota_{\mathfrak{a}}(\alpha)$, as defined in Definition 5. Assuming $|\operatorname{disc}(\mathcal{O})| \leq 2^{2f}$ with $2^{f}|p+1$, we can always find α of norm bounded by 2^{2f} and it suffices to evaluate $\iota_{\mathfrak{a}}(\alpha)$ on a basis of $E_{\mathfrak{a}}[2^{f}]$ to efficiently represent this endomorphism, *e.g.* using the higher dimensional interpolation algorithms from [55, Theorem 3.7 & Section 6.4]. Besides, if $e+2 \leq f$, then knowing $\iota(\alpha)$ on the 2^{f} -torsion is sufficient to compute the 2^{e+2} -torsion lying above ker(F) given by Eq. (5) so to compute the action of any ideal on E. The same holds for $E_{\mathfrak{a}}$: the knowledge of $\iota_{\mathfrak{a}}(\alpha)$ on $E_{\mathfrak{a}}[2^{f}]$ is sufficient to propagate the ideal class group action further.

Now we explain how we compute $\iota_{\mathfrak{a}}(\alpha)$ on a basis of $E_{\mathfrak{a}}[2^f]$. Using the fact that $\bar{\mathfrak{cb}} = \sigma \mathcal{O}$, we obtain that

$$\iota_{\mathfrak{a}}(\alpha) = (\varphi_{\mathfrak{a}})_{*}(\iota) = \frac{1}{\mathrm{N}(\mathfrak{a})}\varphi_{\mathfrak{a}} \circ \iota(\alpha) \circ \widehat{\varphi}_{\mathfrak{a}} = \frac{1}{\mathrm{N}(\mathfrak{b})}\varphi_{\mathfrak{b}} \circ \iota(\alpha) \circ \widehat{\varphi}_{\mathfrak{b}}.$$

Let (P_1, Q_1) and $(P_{\mathfrak{a}}, Q_{\mathfrak{a}})$ be two basis of $E_1[2^f]$ and $E_{\mathfrak{a}}[2^f]$ respectively. Then by Eq. (4), we have $F(\Phi_u(P_1, 0), 0) = ([u]\varphi_{\mathfrak{b}_k}(P_1), 0, *)$ so we can evaluate $[u]\varphi_{\mathfrak{b}_k}(P_1)$, hence $\varphi_{\mathfrak{b}_k}(P_1)$ by multiplying by an inverse of $u \mod 2^f$. We compute $\varphi_{\mathfrak{b}_k}(Q_1)$ similarly. Now, with (easy) discrete logarithm computations in $E_{\mathfrak{a}}[2^f]$, we may find $a, b, c, d \in \mathbb{Z}$ such that $\varphi_{\mathfrak{b}_k}(P_1) = [a]P_{\mathfrak{a}} + [b]Q_{\mathfrak{a}}$ and $\varphi_{\mathfrak{b}_k}(Q_1) =$ $[c]P_{\mathfrak{a}} + [d]Q_{\mathfrak{a}}$. Then, a simple matrix inversion yields

$$\widehat{\varphi}_{\mathfrak{b}_k}(P_\mathfrak{a}) = [\delta \mathcal{N}(\mathfrak{b}_k)]([d]P_1 - [b]Q_1) \text{ and } \widehat{\varphi}_{\mathfrak{b}_k}(Q_\mathfrak{a}) = [\delta \mathcal{N}(\mathfrak{b}_k)](-[c]P_1 + [a]Q_1),$$

with δ an inverse of $ad-bc \mod 2^f$ (which does exist because $N(\mathfrak{b}_k) = \deg(\varphi_{\mathfrak{b}_k})$ is odd, so $\varphi_{\mathfrak{b}_k}$ maps a 2^f -torsion basis to a 2^f -torsion basis). So we have computed $(\widehat{\varphi}_{\mathfrak{b}_k}(P_{\mathfrak{a}}), \widehat{\varphi}_{\mathfrak{b}_k}(Q_{\mathfrak{a}}))$. Similarly, knowing $\varphi_{\mathfrak{b}_e}$, we can compute the action of its dual $\widehat{\varphi}_{\mathfrak{b}_e}$ on the 2^f -torsion provided that $N(\mathfrak{b}_e)$ is odd. We can impose this condition by removing 2 from the set of allowed primes \mathfrak{B} . Hence, we can evaluate $\widehat{\varphi}_{\mathfrak{b}_e} \circ \widehat{\varphi}_{\mathfrak{b}_k} = \widehat{\varphi}_{\mathfrak{b}}$ on the basis $(P_{\mathfrak{a}}, Q_{\mathfrak{a}})$. We can then evaluate $[N(\mathfrak{b})]\iota_{\mathfrak{a}}(\alpha) = \varphi_{\mathfrak{b}} \circ$ $\iota(\alpha) \circ \widehat{\varphi}_{\mathfrak{b}} = \varphi_{\mathfrak{b}_k} \circ \varphi_{\mathfrak{b}_e} \circ \iota(\alpha) \circ \widehat{\varphi}_{\mathfrak{b}_e} \circ \widehat{\varphi}_{\mathfrak{b}_k}$ on $(P_{\mathfrak{a}}, Q_{\mathfrak{a}})$, using our knowledge of the action of $\iota(\alpha)$ on the 2^f -torsion, of $\varphi_{\mathfrak{b}_e}$ and of $\varphi_{\mathfrak{b}_k}$ via F. Since $N(\mathfrak{b}) = N(\mathfrak{b}_e)N(\mathfrak{b}_k)$ is odd, we can invert it modulo 2^f and finally obtain $(\iota_{\mathfrak{a}}(\alpha)(P_{\mathfrak{a}}), \iota_{\mathfrak{a}}(\alpha)(Q_{\mathfrak{a}}))$, as desired.

4 Computing 2-dimensional isogenies of prescribed polarised degree

In this section, we explain how to construct dimension 2 isogenies Φ_u, Φ_v of prescribed polarised degree u, v, in order to solve Eq. (2) (or more precisely the version in Eq. (3)).

We first treat an easy special case in Section 4.1. Then we explain in Section 4.2 how to treat the case of u (or v) an arbitrary positive integer. The drawback of the latter approach is that it involves computing a 4-dimensional 2^e -isogeny to build Φ_u in dimension 2.

4.1 The simplest case: when the polarised degree is \mathfrak{B} -good

Let us first assume that u and v are a sum of two squares. Then, given a curve E and an integer $u = x_u^2 + y_u^2$, the endomorphism

$$M_u = \begin{pmatrix} x_u & -y_u \\ y_u & x_u \end{pmatrix} \tag{8}$$

on E^2 is such that $M_u \widetilde{M}_u = [u]$ is the multiplication by u on each component. Hence M_u induces an isogeny Φ_u of polarised degree u as desired.

The probability that a random number u can be written as sum of two squares is roughly $1/\sqrt{\log(u)} \approx 1/\sqrt{e/2}$. Unfortunately, writing u as above requires the factorisation of u, so instead we look for u a prime congruent to 1 modulo 4, which reduces the probability to $1/\log(u) \approx 1/e$. Since we need both u and v to be of this form, we expect to find a valid pair after $\approx e^2$ attempts. Looking at the size of the primes we are working with, this approach become soon too expensive.

Instead, we can adopt the same strategy as in Section 3.2 to relax our condition and aim for values \mathfrak{B} -good values u, v, for a given set of small primes.

Definition 6. Let \mathfrak{B} be a set of small primes. We say that $u \in \mathbb{N}^*$ is \mathfrak{B} -good when u is of the form $u = g_u(x_u^2 + y_u^2)$ where $x_u, y_u \in \mathbb{N}$ and $g_u \in \mathbb{N}^*$ is a product of primes in \mathfrak{B} .

We may then write $u = g_u(x_u^2 + y_u^2)$ and $v = g_v(x_v^2 + y_v^2)$, as above, where g_u and g_v contain all the prime factors in \mathfrak{B} . In this case, we can restrict to primes that are 3 mod 4, since the ones that are 1 mod 4 can already be expressed as sum of squares, and so can their products. Notice that small primes are more likely to appear in the factorization of a number, and consequently small factors that are 3 mod 4 are more likely to prevent a number from being a sum of two squares. Hence, removing them significantly increase the probability of finding a solution. If we choose \mathcal{O} such that these small primes (e.g. 3, 7, 11) are split in \mathcal{O} , we can easily compute isogenies of degrees g_u and g_v via the action of $\mathrm{Cl}(\mathcal{O})$. In particular, in the case of CSIDH, these isogenies are defined over \mathbb{F}_p so they are even more efficient to compute (see Section 5.2).

Given a pair u, v with no factors in \mathfrak{B} , we are now left with the task of determining if both numbers can be expressed as sums of two squares. It is well-known that an integer can be written as a sum of two squares if and only if, in its prime power decomposition, there is no prime power ℓ^k such that k is odd and $\ell \equiv 3 \mod 4$. Unfortunately, as mentioned above, checking this condition requires factoring, which we cannot afford. What we can do instead is to perform trial division up to a fixed bound, and then test what is left for primality. If the primality test succeeds, and all nonsquare factors are 1 mod 4, we can apply Cornacchia's algorithm to obtain x_u and y_u (respectively x_v, y_v). This strategy is further discussed in Section 5.2 in the case of CSIDH.

Remark 4.1. It is possible to relax the \mathfrak{B} -good condition on (u, v) by allowing u (resp. v) to take the form $g_u(ax_u^2 + by_u^2)$ (resp. $g_v(ax_v^2 + by_v^2)$), where $g_u, g_v, a, b \in \mathfrak{B}$, while ensuring that Kani's lemma remains applicable as in Section 3.1. Then, at the cost of computing 4 additional isogenies of degree a and b with Elkies' algorithm, we improve the likelihood that u and v are suitable for computing isogenies of degree u and v in dimension 2. Experiments have shown that allowing a and b to take values in $\{2, 3, 5, 7\}$ increases the probability of success by approximately a factor of 3. Further details on this construction can be found in Appendix C.

4.2 The general case

We now treat the case of a general u, v. Although it is easy to construct Φ_u, Φ_v in dimension 4, we saw in Section 3.1 that this would involve a dimension 8 isogeny for F. In this section we explain how to construct them in dimension 2,

on E^2 . For that, we extend the approach of QFESTA [44] from dimension 1 to dimension 2. A drawback is that our approach will be heuristic, although our implementation shows that the heuristic works well in practice.

Let (E, ι) be an \mathcal{O} -oriented supersingular elliptic curve. Let $\Delta = \operatorname{disc}(\mathcal{O})$ be the discriminant of \mathcal{O} . First, we show that we can heuristically build (efficient) endomorphisms on E^2 of polarised degree N, as long as $N \gg |\Delta|$. By assumption, we can evaluate any endomorphism $\gamma \in \mathcal{O}$, which gives us a 1-dimensional isogeny $\iota(\gamma)$ of degree $N(\gamma)$. Since $\sqrt{\Delta} \in \mathcal{O}$, the quadratic form $\gamma \mapsto N(\gamma)$ restricts on the suborder $\mathbb{Z} + \sqrt{\Delta}\mathbb{Z}$ of \mathcal{O} to the quadratic form $q(x, y) = x^2 + |\Delta|y^2$.

Now let $\gamma_1, \gamma_2 \in \mathcal{O}$ of norms n_1, n_2 respectively, and consider the endomorphism

$$\gamma = \begin{pmatrix} \iota(\gamma_1) & \iota(\overline{\gamma_2}) \\ -\iota(\gamma_2) & \iota(\overline{\gamma_1}) \end{pmatrix} \in \operatorname{End}(E^2).$$

Then, by [55, Lemma 3.2], its polarised dual is

$$\widetilde{\gamma} = \begin{pmatrix} \iota(\overline{\gamma_1}) & -\iota(\overline{\gamma_2}) \\ \iota(\gamma_2) & \iota(\gamma_1) \end{pmatrix},$$

so that $\tilde{\gamma} \circ \gamma = [n_1 + n_2]$ in End (E^2) and γ is an $(n_1 + n_2)$ -isogeny. In particular, we can build endomorphisms on E^2 of polarised degree N as long as N is represented by the quadratic form $q(x, y, z, t) = (x^2 + y^2) + |\Delta|(z^2 + t^2)$.

Heuristic 7. Let $N \gg |\Delta|$. Then we can heuristically build an efficient endomorphism $\gamma \in \text{End}(E^2)$ of polarised degree N.

Justification. Since $N \gg |\Delta|$, we can sample many small z, t such that $(z^2 + t^2)|\Delta| < N$. Then if $m = N - |\Delta|(z^2 + t^2)$ is a sum of two squares, we can find γ by the discussion above. This situation can be efficiently detected whenever m is a prime congruent to 1, which heuristically happens with probability $\approx 1/\log(N)$. Since $N \gg |\Delta|$, we can try with many different couples (z, t) until one succeeds.

In practice, we can use a direct adaptation of the algorithm RepresentInteger from [23] to find a solution of $(x^2 + y^2) + |\Delta|(z^2 + t^2) = N$.

Now by Lemma 3, if γ is an efficiently represented endomorphism of E^2 of polarised degree $N = N_1 N_2$, with N_1, N_2 coprime, then it decomposes uniquely as $\gamma = \mu_2 \circ \gamma_1 = \mu_1 \circ \gamma_2$ where the γ_i, μ_i are isogenies of polarised degree N_i . Furthermore, by Kani's lemma (Lemma 4) the 4-dimensional isogeny

$$\Gamma = \begin{pmatrix} \gamma_1 & \widetilde{\mu_2} \\ -\gamma_2 & \widetilde{\mu_1} \end{pmatrix} \tag{9}$$

is of polarised degree $N_1 + N_2$. The kernel of Γ is given by Ker $\Gamma = \{(N_1P, \gamma P) \mid P \in E^g[N_1 + N_2]\}$, so we can efficiently evaluate Γ to obtain the 2-dimensional isogeny γ_1 of polarised degree N_1 as long as the $(N_1 + N_2)$ -torsion on E is smooth and accessible (*i.e.* defined over a small extension of \mathbb{F}_p). Hence, an efficient representation of γ yields an efficient representation of γ_1 in the sense of Definition 5.

For simplicity, and because this matches our implementation choice, we will now restrict to the case where we look for $N_1 + N_2 = 2^e$ a power of two. **Theorem 8.** Assume that $E[2^e] \subseteq E(\mathbb{F}_{p^2})$ (with $e \geq 3$), that $2^{3e/2-1} \gg |\Delta|$, and that $u < 2^{e-1}$ is odd. Then, under Heuristic 7, we can efficiently represent an isogeny Φ_u of polarised degree u on E^2 .

Proof. First consider $N = N_1 N_2$ with $N_1 = u, N_2 = 2^e - u$. Since u is assumed to be odd, N_1 is coprime to N_2 . If $N \gg |\Delta|$, we can use Heuristic 7 to build an endomorphism in dimension 2 of polarised degree N, which we split via a 4-dimensional isogeny of polarised degree 2^e to get Φ_u .

If this is not the case, then since $2^{3e/2} \gg |\Delta|$ by assumption, this means that u is smaller than 2^f , with $f = \lceil e/2 \rceil$. We use a two step solution like in [45, 27]. First we let $u_2 = u(2^f - u)$, then $2^f - 1 \le u_2 < 2^{2f-2} \le 2^{e-1}$. Then we let $N = u_2(2^e - u_2)$. By our choice of $f, N \ge 2^{3e/2-1}$, so $N \gg |\Delta|$ and we can use Heuristic 7 to build an endomorphism in dimension 2 of polarised degree N, which we split a first time (via a 4-dimensional isogeny of polarised degree 2^e) to get a 2-dimensional isogeny Φ_{u_2} of polarised degree u_2 , which we split again (via a 4-dimensional isogeny of polarised degree 2^f) to get Φ_u .

Remark 9.

- To compute Γ as defined in Eq. (9), we only need the *m*-torsion to be smooth and accessible on *E*, where $N_1 + N_2 \mid m^2$, see [57, § 6.3 and Appendix B]. This allows to relax the conditions on Theorem 8 to $u < 2^{2e}$ and $2^{3e-1} \gg \Delta$.
- In practice, for our application from Section 3, we have $u, v \approx \sqrt{|\Delta|}$. Furthermore, in the case of CSIDH, we have $\Delta = -p$, and we select p such that $p = c2^f 1$ for a small cofactor c. In particular, we can take e = f in Theorem 8, so that $2^e \approx p$. In that case we already have $u(2^e u) \gg p$, and we do not need the two step solution, nor the more relaxed torsion condition from the item above.

Combining Section 3 with Theorem 8, the resulting algorithm to compute the action of a quadratic ideal $\mathfrak{a} \subset \mathcal{O}$ on an \mathcal{O} -oriented elliptic curve is very similar to the algorithm used in [5] to convert a quaternionic ideal to an isogeny when the full endomorphism ring of E is known. We make up the lack of known endomorphisms on E to build isogenies of suitable fixed degree by going up to dimension 2 and working on E^2 . This doubling of dimension propagates to the rest of the algorithm, and we end up computing Φ_u, Φ_v and $E_{\mathfrak{a}}$ via an isogeny of dimension 4, compared to dimension 2 in [5].

This change of dimension has important consequences for the fine tuning of the algorithm. Indeed, as we have seen in Section 4.1, Φ_u (resp. Φ_v) are much easier to compute whenever u (resp. v) is a sum of two squares, or at least \mathfrak{B} good in the sense of Definition 6. In particular, in that case we do not need to use a 4-dimensional isogeny to compute Φ_u .

A similar situation happened in [5]: if u was a sum of two squares we could compute Φ_u directly without going through a 2-dimensional isogeny. However, experiments showed that spending more time on the norm equation to find ua sum of two squares did not compensate having to do one less 2-dimensional isogeny. The situation is different in our case, because a 4-dimensional 2^e -isogeny is much slower than a 2-dimensional 2^e -isogeny (by a factor roughly $\times 8$). Depending on the size of our parameters, experiments show that the optimal choices depend on the size of Δ .

- If Δ is of moderate size, we look for \mathfrak{B} -good u, v, and only compute one 4-dimensional isogeny for $E_{\mathfrak{a}}$.
- If Δ is of large size, we look for only one of u, v to be \mathfrak{B} -good, while we let the other one be arbitrary. This time computing $\varphi_{\mathfrak{a}}$ requires two 4-dimensional isogenies. For CSIDH, our experiments in Section 6.4 suggest this strategy becomes faster than the previous one for p of 4000 bits. We remark that for very large Δ , we expect that letting u, v be arbitrary and computing three 4-dimensional isogenies would be the optimal choice.

In the case of CSIDH (detailed in Sections 5 and 6), only the first choice is implemented and the computation stays reasonably efficient even for our biggest parameters (p of 4000 bits) as long as the allowed set of small split primes \mathfrak{B} is well chosen. We refer to Section 6.4 for further discussion on the above trade-off.

5 Our algorithm specialized to CSIDH

In this section, we apply our algorithm to the CSIDH setting, or more accurately, the CSURF setting [13]; we fix a prime of the form $p = c2^f - 1$ where c is a small odd integer and $f \gg 1$ so that $p \equiv 7 \pmod{8}$, and work with curves oriented by $\mathbb{Z}\left[\frac{1+\sqrt{-p}}{2}\right]$, where the orientation is given by sending $\sqrt{-p}$ to the Frobenius endomorphism. Not only does using the orientation given by Frobenius make evaluating the orientation very fast, but by using the maximal order instead of the typical CSIDH suborder $\mathbb{Z}[\sqrt{-p}]$, we are able to execute our whole algorithm over \mathbb{F}_p , by choosing a particularily suitable basis of $E[2^e]$, and working with x-only arithmetic. This is further detailed in Section 5.2 and Appendix D.

We now detail how to evaluate the action of any primitive ideal $\mathfrak{a} \subset \mathbb{Z}\left[\frac{1+\sqrt{-p}}{2}\right]$.

5.1 Step 1: the norm equation

In this section, we explain how to solve the norm equation (3) sketched in Section 3.2. As in Section 4.1, we fix a set of small Elkies primes \mathfrak{B} , *i.e.* of primes that split in $\mathbb{Q}(\sqrt{-p})$ (including 2). We begin by creating a list of primitive ideals equivalent to \mathfrak{a} . More precisely, we obtain a list of equivalent ideals $\mathfrak{a}^{(i)}$ such that $\mathfrak{a}^{(i)} = \mathfrak{a}_e^{(i)}\mathfrak{a}_k^{(i)}$ and $N(\mathfrak{a}_e^{(i)})$ is a product of primes in \mathfrak{B} . We then want to find a pair (i, j) such that for $N_1 = N(\mathfrak{a}_k^{(i)})$ and $N_2 = N(\mathfrak{a}_k^{(j)})$ (note that the smooth part has been removed from the norm), we can find positive integers u, v solving the equation

$$uN_1 + vN_2 = 2^e. (10)$$

Since generically N_1N_2 determines the minimal value for e such that the above equation has a solution, we test pairs in order of increasing N_1N_2 .

Given such pair of norms N_1, N_2 , we first check that they are coprime. Indeed, if $gcd(N_1, N_2)$ is not a power of 2, Equation (10) has no solutions. Since $2 \in \mathfrak{B}$ and we assume N_1, N_2 has no prime factors in \mathfrak{B} , this requirement simply becomes $gcd(N_1, N_2) = 1$.

If $gcd(N_1, N_2) = 1$, we can find u_0 and v_0 such that

$$u_0 N_1 + v_0 N_2 = 1$$

via the extended Euclidean algorithm, and consequently

$$2^e u_0 N_1 + 2^e v_0 N_2 = 2^e.$$

Note that one of u_0 and v_0 will be negative and that we parametrize all possible solutions to the equation by writing

$$(2^e u_0 + kN_2)N_1 + (2^e v_0 - kN_1)N_2 = 2^e$$

for some $k \in \mathbb{Z}$. This also allows to determine the minimal e for which Equation (10) has a solution by enumerating the set of k for which both $u = 2^e u_0 + kN_2$ and $v = 2^e v_0 - kN_1$ are positive.

Remark 5.1. The exponent e in Equation (10) determines the number of fourdimensional 2-isogeny steps to be computed, so taking a lower exponent e will be more efficient.

As remarked before and also observed in [6], solving Equation (10) directly for $\mathfrak{a}^{(i)}$ and $\mathfrak{a}^{(j)}$, often fails since N_1, N_2 are in $O(\sqrt{p})$ and $2^e < p$, which makes the original Clapoti approach unpractical in our case. Removing factors in \mathfrak{B} , makes N_1 and N_2 much smaller and hence greatly increases the probability of finding solutions to Equation (10): the number of expected solutions grows roughly as the product of all factors removed. This leads to the following tradeoff: the bigger the set of primes \mathfrak{B} , the higher the probability of finding a solution, but the more time is spent in the second step computing small degree isogenies. Furthermore, even if the probability of finding a solution, we can simply rerandomize the starting ideal (by some small ideal) and start over. More details about this approach are given below.

Assuming that we can find enough solution pairs (u, v), for each one we have to determine whether u and v are \mathfrak{B} -good, *i.e.* if we can write $u = g_u(x_u^2 + y_u^2)$ and $v = g_v(x_v^2 + y_v^2)$ with the prime factors of g_u and g_v in \mathfrak{B} . Recall that a number can be expressed as a sum of two squares if and only if all its prime factors that are 3 mod 4 appear with an even exponent. The obvious choice is then to set g_u (resp. g_v) to be the product of all prime factors that are 3 mod 4 and contained in \mathfrak{B} and dividing u (resp. v). To determine whether what is left contains more prime factors 3 mod 4 we would need to factor u, which is too expensive in general. As such we simply proceed by trial division up to a small bound, hoping that what is left at the end is a prime that is 1 mod 4 (note that we simply reject when the left over is 3 mod 4). If everything succeeds, we have found the factorization of u, and hence its decomposition as sum of two squares. The same procedure applies to v. How to choose an optimal set \mathfrak{B} is analyzed in greater detail in Section 6.1.

Once we have found a full solution, we proceed to the next step: we will denote the ideals $\mathfrak{a}^{(i)}, \mathfrak{a}^{(j)}$ as \mathfrak{b} and \mathfrak{c} , so in particular, $N_1 = N(\mathfrak{b}_k)$ and $N_2 = N(\mathfrak{c}_k)$.

Rerandomization For some ideals, Equation (10) may not have a solution with u, v of the required shape. As already observed, increasing the bound \mathfrak{B} is always a possible solution, though this quickly becomes expensive.

Another solution to this is to simply rerandomize the starting ideal \mathfrak{a} . Namely, if none of the sampled, equivalent ideals gives us a solution for Equation (10), we can multiply \mathfrak{a} by a non-principal ideal \mathfrak{l} for which the action is easy to evaluate (*e.g.* an ideal above ℓ , where ℓ is the smallest split prime). We obtain a new ideal $\mathfrak{a}' = \mathfrak{a}\mathfrak{l}$ and try to solve Equation (10) for this ideal instead. If we manage to do so, we can simply run our algorithm on \mathfrak{a}' , using this solution, starting from the starting curve $E_{\overline{\mathfrak{l}}} = \overline{\mathfrak{l}} \cdot E$ instead, which is easy to compute by the choice of \mathfrak{l} . Otherwise, we can rerandomize again by computing $\mathfrak{a}'' = \mathfrak{l}\mathfrak{a}'$, and so on. As soon as Equation (10) has a reasonable chance of being solved, this method takes care of failures quite efficiently, making Step 1 faster at the cost of a few Elkies steps.

5.2 Step 2: computing kernel points

After determining a suitable pair of ideals $\mathfrak{b}, \mathfrak{c}$, our goal is to construct the 4dimensional 2^e -isogeny described in Section 3.2. Let E denote the starting curve on which we want to act with \mathfrak{a} , then we obtain the following diagram derived from the isogeny diamond in Section 3.2:



In the diagram above, we have:

 $- \Psi_1 = \operatorname{diag}(\varphi_{\mathfrak{b}_e}, \varphi_{\mathfrak{b}_e})$, where $\varphi_{\mathfrak{b}_e}$ is the isogeny corresponding to the action of \mathfrak{b}_e ; analogously, $\Psi_2 = \operatorname{diag}(\varphi_{\mathfrak{c}_e}, \varphi_{\mathfrak{c}_e})$;

The Clapoti Cult.

- $-\Phi_1 = \operatorname{diag}(\varphi_{\mathfrak{b}_k}, \varphi_{\mathfrak{b}_k}) \text{ and } \Phi_2 = \operatorname{diag}(\varphi_{\mathfrak{c}_k}, \varphi_{\mathfrak{c}_k});$
- Φ_u and Φ_v are isogenies of polarised degree u and v respectively; as explained in Section 4.1, we can write $\Phi_u = M_u \operatorname{diag}(\varphi_u, \varphi_u)$, with $\operatorname{deg}(\varphi_u) = g_u, \Phi_v =$ $M_v \operatorname{diag}(\varphi_v, \varphi_v)$ with $\operatorname{deg}(\varphi_v) = g_v, M_u$ and M_v being given by Equation (8).

Further, the isogenies Φ and Ψ are the isogenies of polarised degree uN_1 and vN_2 completing the square, whose existence is guaranteed by Lemma 3. In this setting we can also describe them explicitly, as the following lemma shows.

Lemma 5.1. Φ is of the form $\Phi'_1 \circ \widetilde{\Phi}'_u$, where:

- $\Phi'_{1} := \operatorname{diag}(\varphi'_{\mathfrak{b}_{k}}, \varphi'_{\mathfrak{b}_{k}}) : E'_{1}^{2} \to E^{2}_{v} \text{ with } E'_{1} := [\overline{\mathfrak{b}}_{k}] \cdot E_{v} \text{ and } \varphi'_{\mathfrak{b}_{k}}, \text{ the isogeny associated to the action of } \mathfrak{b}_{k} \text{ on } E'_{1}; \\ \Phi'_{u} := M_{u} \operatorname{diag}(\varphi'_{u}, \varphi'_{u}) : E'_{1}^{2} \to E'^{2}, M_{u} \text{ being given by Equation (8) and } \varphi'_{u} \text{ being the isogen of degree } g_{u} \text{ given by the action of the same ideal } I_{u} \text{ as } f(\overline{\mathfrak{b}}_{k}) \in \mathbb{R}^{d}$ φ_u (as a product of Elkies primes).

 Ψ is of the form $\Phi'_{v} \circ \widetilde{\Phi}'_{2}$, where:

- $\Phi'_{2} := \operatorname{diag}(\varphi'_{\mathfrak{c}_{k}}, \varphi'_{\mathfrak{c}_{k}}) : E'_{2}^{2} \to E^{2}_{u} \text{ with } E'_{2} := [\overline{\mathfrak{c}}_{k}] \cdot E_{u} \text{ and } \varphi'_{\mathfrak{c}_{k}}, \text{ the isogeny associated to the action of } \mathfrak{c}_{k} \text{ on } E'_{2};$ $\Phi'_{v} := M_{v} \operatorname{diag}(\varphi'_{v}, \varphi'_{v}) : E'_{2}^{2} \to E'^{2}, M_{v} \text{ being given by Equation (8) and } \varphi'_{v} \text{ being the isogeny of degree } g_{v} \text{ given by the action of the same ideal } I_{v} \text{ as } \varphi_{v}$ (as a product of Elkies primes).

In particlar, the common codomain of $\widetilde{\Phi}$ and Ψ is a product of elliptic curves E'^2 .

Proof. We have to verify that $\Phi'_1 \circ \tilde{\Phi}'_u$ and $\Phi'_v \circ \tilde{\Phi}'_2$ defined above are $N_1 u$ and N_2v -isogenies respectively making the diagram commute, *i.e.* such that:

$$\Phi_1' \circ \tilde{\Phi}_u' \circ \Phi_v' \circ \tilde{\Phi}_2' = \Phi_v \circ \tilde{\Phi}_2 \circ \Phi_1 \circ \tilde{\Phi}_u.$$
⁽¹¹⁾

First, we verify that the composition on the left makes sense *i.e.* that Φ'_u and \varPhi'_v have the same codomain. By definition, the codomain of \varPhi'_u is

$$E' = [I_u]E'_1 = [I_u\overline{\mathfrak{b}}_k]E_v = [I_u\overline{\mathfrak{b}}_kI_v]E_2 = [I_uI_v\overline{\mathfrak{b}}_k\mathfrak{c}_e]E$$

and the codomain of Φ'_v is

$$E'' = [I_v]E'_2 = [I_v\bar{\mathfrak{c}}_k]E_u = [I_v\bar{\mathfrak{c}}_kI_u]E_1 = [I_uI_v\bar{\mathfrak{c}}_k\mathfrak{b}_e]E.$$

But $\mathfrak{b} = \mathfrak{b}_e \mathfrak{b}_k$ and $\mathfrak{c} = \mathfrak{c}_e \mathfrak{c}_k$ are both equivalent to \mathfrak{a} . It follows that $\overline{\mathfrak{b}}_k \mathfrak{c}_e$ and $\overline{\mathfrak{c}}_k \mathfrak{b}_e$ are equivalent, so that $E' \simeq E''$.

By construction, $\Phi'_1 \circ \tilde{\Phi}'_u$ and $\Phi'_v \circ \tilde{\Phi}'_2$ are $N_1 u$ and $N_2 v$ -isogenies respectively, so we only have to prove Equation (11). On the one hand, we have:

$$\varPhi_1' \circ \widetilde{\varPhi}_u' \circ \varPhi_v' \circ \widetilde{\varPhi}_2' = \widetilde{M}_u \circ M_v \circ \operatorname{diag}(\varphi_{\mathfrak{b}_k}' \circ \widehat{\varphi}_u' \circ \varphi_v' \circ \widehat{\varphi}_{\mathfrak{c}_k}', \varphi_{\mathfrak{b}_k}' \circ \widehat{\varphi}_u' \circ \varphi_v' \circ \widehat{\varphi}_{\mathfrak{c}_k}'),$$

since M_u and M_v commute with diagonal isogenies. On the other hand:

$$\Phi_v \circ \Phi_2 \circ \Phi_1 \circ \Phi_u = M_v \circ M_u \circ \operatorname{diag}(\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k} \circ \widehat{\varphi}_u, \varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k} \circ \widehat{\varphi}_u).$$

Both $\varphi'_{\mathfrak{b}_k} \circ \widehat{\varphi}'_u \circ \varphi'_v \circ \widehat{\varphi}'_{\mathfrak{c}_k}$ and $\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k} \circ \widehat{\varphi}_u$ correspond to the action of the ideal $\overline{I}_u I_v \overline{\mathfrak{c}}_k \mathfrak{b}_k$ on E_u so these isogenies must be equal. Besides, a simple matrix computation ensures that \widetilde{M}_u and M_v commute. This proves Equation (11) and the lemma.

Let (P, Q) be a basis of $E_1[2^{e+2}]$ (recall from Section 3.1 that the theta isogeny algorithm requires the 2^{e+2} -torsion to compute a 2^e -isogeny, see Appendix A for more details). To compute the kernel of the isogeny in dimension 4, we need to compute the points

$$P_u = \varphi_u(P), \quad Q_u = \varphi_u(Q)$$

on E_u and

$$P_v = \varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(P), \quad Q_v = \varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(Q),$$

on E_v .

Isogenies of degree g_u and g_v The isogenies φ_u and φ_v with $\deg(\varphi_u) = g_u$ and $\deg(\varphi_v) = g_v$ have no specific requirement other than their degree. Although it is possible to deal with all small primes via Vélu's algorithm [62], it is more efficient to restrict \mathfrak{B} to Elkies primes (*i.e.* splitting primes in $\mathbb{Q}(\sqrt{-p})$) and we can use the corresponding algorithm [29] to compute the small degree isogenies.

Note that φ_u of degree g_u and φ_v of degree g_v can be used to construct 2-dimensional isogenies of polarised degree $u = g_u(x_u^2 + y_u^2)$ and $v = g_v(x_v^2 + y_v^2)$ respectively by composition with matrices M_u, M_v described in Section 4.1. However, we do not need to perform that step now, as it can be included in the 4-dimensional computation.

Working over \mathbb{F}_p Notice that, by definition, the isogenies $\varphi_{\mathfrak{b}}$ and $\varphi_{\mathfrak{c}}$ are \mathbb{F}_p rational, and as noted above, our choice of primes in \mathfrak{B} imply that φ_u and φ_v can
also be chosen to be defined over \mathbb{F}_p . The main difficulity comes from evaluating
these isogenies on a basis of $E[2^{e+2}]$, which is not defined over \mathbb{F}_p .

We will show that all operations can infact be carried out over the Kummer line, where we have a rational basis. Let f the largest integer so that $2^{f} \mid (p+1)$. By our choice of prime, the two eigenvalues of Frobenius on $E[2^{f-1}]$ are ± 1 . Thus, we can pick a basis $\langle P, Q \rangle = E[2^{f-1}]$ corresponding to the two eigenvectors of Frobenius; we let P, Q be points of order 2^{f-1} satisfying

$$\pi(P) = P, \quad \pi(Q) = -Q. \tag{12}$$

Clearly, these points P, Q are \mathbb{F}_p -rational on the kummer line $E/\pm 1$, and together (adjusting by translation by a point of 2-torsion if needed) they form a basis of

 $E[2^{f-1}]$. This is where we use that we are on the surface. If we were on the floor, picking similar points would not give a basis, since in that case, $[2^{f-2}]P = [2^{f-2}]Q$, the unique rational point of 2-torsion.

In Appendix D.1 we show how to efficiently generate such a basis, based on a technique which is similar to how one usually computes bases of $E[2^f]$ over \mathbb{F}_{p^2} .

Evaluating the ideals The rest of this section will be devoted to the computation of P_v, Q_v . In particular, we need to evaluate $\widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}$ on a basis of $E_1[2^{e+2}]$. From Equation (6) we know that

$$\widehat{\varphi}_{\mathfrak{c}_{k}} \circ \varphi_{\mathfrak{b}_{k}} = \frac{1}{N(\mathfrak{b}_{e})N(\mathfrak{c}_{e})}\varphi_{\mathfrak{c}_{e}} \circ \iota(\sigma) \circ \widehat{\varphi}_{\mathfrak{b}_{e}}$$
(13)

with $\bar{\mathfrak{c}}\mathfrak{b} = \sigma \mathcal{O}$. By construction, $N(\mathfrak{b}_e)$ and $N(\mathfrak{c}_e)$ are both smooth, with all their factors in \mathfrak{B} . Let us assume for simplicity that they are also odd; the even case needs a bit more care and is detailed afterwards.

We detail how to evaluate the endomorphism $\iota(\sigma)$. First, assume that σ is of the form $a+b\sqrt{-p}$, with $a, b \in \mathbb{Z}$. Explicitly, this is mapped to the endomorphism $[a] + [b]\pi$, where π denotes the Frobenius endomorphism. Evaluating such an endomorphism on our choice of basis is particularily easy: with the same notation as in the previous section, the evaluation is simply given by

$$[a]P + [b]\pi(P) = [a]P + [b]P = [a+b]P,$$

$$[a]Q + [b]\pi(Q) = [a]Q + [b](-Q) = [a-b]Q.$$

Notice in particular that we can evaluate this on the Kummer line. Although σ is an element of the order $\mathbb{Z}[\frac{1+\sqrt{-p}}{2}]$, and hence is more generally of the form $a + b\sqrt{-p}$, where a, b are in $\mathbb{Z}[\frac{1}{2}]$, *i.e.* are rational numbers with denominator at most 2, we show in Appendix D.2 how to evaluate this on our basis (on the Kummer line) without having to use division points, which may not be \mathbb{F}_p -rational.

Further, note that since we are after the action on the 2^{e+2} -torsion, we can compute $(N(\mathfrak{b}_e)N(\mathfrak{c}_e))^{-1} \pmod{2^{e+2}}$, to account for the division in Equation (13). Hence, all that remains to evaluate $\widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}$ is to describe how we efficiently recover the isogenies $\varphi_{\mathfrak{b}_e}$ and $\varphi_{\mathfrak{c}_e}$.

Notice that these isogenies are (by definition) the product of prime degree Elkies isogenies. More specifically, say that for a given prime l, we have $v_l(N(\mathfrak{b}_e)) = k$, *i.e.* l divides $N(\mathfrak{b}_e)$ exactely k times. Then \mathfrak{b}_e will act with exactely k degree-l Elkies isogenies. We can also assume that all these isogenies will be in the same direction, since otherwise \mathfrak{b}_e and consequently \mathfrak{b} would factor through multiplication by l, and we assumed that \mathfrak{b} was primitive (if it was not, the ideal \mathfrak{b}/l would also be equivalent to \mathfrak{a} , but with smaller norm).

With Elkies' algorithm, we can return both kernel polynomials of degree l determining the two Elkies isogenies of degree l. To check which polynomial corresponds to the correct isogeny, we check the eigenvalue of Frobenius. In more

detail, assume that $(\ell, \sqrt{-p} - \lambda) = \mathfrak{l} \subseteq \mathfrak{b}_e$, *i.e.* we wish to evaluate the action of \mathfrak{l} . Since the roots of the correct kernel polynomial h(x) are precisely the *x*-coordinates of the points of order ℓ , satisfying $\pi(P) = [\lambda]P$, we simply test this equality symbolically: we have the equality

$$(x^p, y^p) \equiv [\lambda](x, y) \pmod{h(x), y^2 - f(x)},$$

where f(x) is the polynomial coming from the curve equation. If these values are not equal, we know that this kernel polynomial corresponds to the other eigenvalue. Note that when we are doing multiple ℓ -isogenies, we need only check the eigenvalue of Frobenius for the first ℓ -isogeny; after that, we can recognise the right kernel polynomial to be the one not corresponding to the previous *j*-invariant.

Remark 5.2. If \mathfrak{b}_e and \mathfrak{c}_e share some steps, we can perform those steps (corresponding to the action of the ideal $\mathfrak{b}_e + \mathfrak{c}_e$) in advance, and start evaluating the ideal action from the resulting curve. The advantage is twofold: first, we avoid evaluating the same ideal twice. Second, we do not need to evaluate the common part on points. We can instead start computing P_u and P_v directly from the codomain curve.

Even norm part We now explain how to deal with even normed ideals. For this, we write the ideal \mathfrak{b}_e as

$$\mathfrak{b}_e = \mathfrak{b}_{e'}\mathfrak{b}_2$$

where \mathfrak{b}_2 is an ideal of norm $2^{f_{\mathfrak{b}}}$, and correspondingly for \mathfrak{c}_e . As explained in Remark 5.2, the shared steps of \mathfrak{b}_e , \mathfrak{c}_e can be computed at the beginning, hence we can assume that \mathfrak{b}_2 and \mathfrak{c}_2 are coprime (*i.e.* the corresponding 2-isogenies go in "opposite" directions). Writing $f_{12} = f_{\mathfrak{b}} + f_{\mathfrak{c}}$, we get that

$$\widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k} = \frac{1}{N(\mathfrak{b}_{e'})N(\mathfrak{c}_{e'})2^{f_{12}}} \varphi_{\mathfrak{c}_e} \circ \iota(\sigma) \circ \widehat{\varphi}_{\mathfrak{b}_e}.$$

The only added difficulity that need to be accounted for is that we cannot find an inverse to $2^{f_{12}}$ modulo 2^{e+2} .

We fix this by taking a basis of a larger torsion group. Using the same technique as for the odd normed case, we can thus evaluate

$$[2^{f_{12}+1}] \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k} = \frac{1}{N(\mathfrak{b}_{e'})N(\mathfrak{c}_{e'})} \varphi_{\mathfrak{c}_e} \circ \iota(\sigma) \circ \widehat{\varphi}_{\mathfrak{b}_e},$$

and hence we can find the evaluation of $\widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}$ on $E_1[2^{e+2}]$ by instead starting with a basis of $E[2^{e+2+f_{12}}]$. (We remark that by construction, we have $e+2+f_{12} \leq f$.)

5.3 Step 3: computing the 4-dimensional isogeny

In this step, we compute the 4-dimensional 2^e -isogeny isogeny $F : A_u \times A_v \to E_a^2 \times A$ from Section 3.2 (defined in Equation (4)). We can actually be much more explicit than in Section 3.2. From Section 5.2, we obtain that $A_u = E_u^2$, $A_v = E_v^2$ and $A = E'^2$. In particular

$$F = \begin{pmatrix} \Phi_1 \circ \widetilde{\Phi}_u & \Phi_2 \circ \widetilde{\Phi}_v \\ -\Phi'_v \circ \widetilde{\Phi}'_2 & \Phi'_u \circ \widetilde{\Phi}'_1 \end{pmatrix} : E_u^2 \times E_v^2 \to E_a^2 \times {E'}^2.$$

Equation (5) also becomes:

$$\ker(F) = \left\{ ([uN_1]\varphi_u(P), [uN_1]\varphi_u(Q), \\ [g_u]M_v \cdot M_u^T(\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(P), \varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(Q))) \mid P, Q \in E_u[2^e] \right\} \\ = \left\{ ([N_1]M_u(\varphi_u(P), \varphi_u(Q)), \\ M_v(\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(P), \varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(Q))) \mid P, Q \in E_1[2^e] \right\}.$$
(14)

To compute F, we proceed with theta coordinates of level 2 and the algorithms from [18] (see Appendix B). These algorithms require a basis of an isotropic subgroup (for the Weil pairing) $K' \subset (E_u^2 \times E_v^2)[2^{e+2}]$ such that $[4]K' = \ker(F)$.

Let (P,Q) be a basis of $E_1[2^{e+2}]$, $\zeta := e_{2^{e+2}}(P,Q)$, η_1, α be inverses of N_1, uN_1 modulo 2^{e+2} respectively, $(P_u, Q_u) := \varphi_u(P,Q)$ and $(P_v, Q_v) := \varphi_v \circ \widehat{\varphi}_{\mathfrak{e}_k} \circ \varphi_{\mathfrak{b}_k}(P,Q)$ the points computed in Step 2. Then, applying Lemma 12.(i) to F and the ζ -symplectic basis $(x_1, x_2, y_1, y_2) := ((\varphi_{\mathfrak{b}_k}(P), 0), (0, \varphi_{\mathfrak{b}_k}(P)), ([\eta_1]\varphi_{\mathfrak{b}_k}(Q), 0), (0, [\eta_1]\varphi_{\mathfrak{b}_k}(Q)))$, we obtain a basis

$$T_{1} := ([N_{1}x_{u}]P_{u}, [N_{1}y_{u}]P_{u}, [x_{v}]P_{v}, [y_{v}]P_{v})$$

$$T_{2} := ([-N_{1}y_{u}]P_{u}, [N_{1}x_{u}]P_{u}, [-y_{v}]P_{v}, [x_{v}]P_{v})$$

$$T_{3} := ([(1 - 2^{e}\alpha)x_{u}]Q_{u}, [(1 - 2^{e}\alpha)y_{u}]Q_{u}, [\eta_{1}x_{v}]Q_{v}, [\eta_{1}y_{v}]Q_{v})$$

$$T_{4} := ([-(1 - 2^{e}\alpha)y_{u}]Q_{u}, [(1 - 2^{e}\alpha)x_{u}]Q_{u}, [-\eta_{1}y_{v}]Q_{v}, [\eta_{1}x_{v}]Q_{v}),$$

of an isotropic subgroup $K' \subset (E_u^2 \times E_v^2)[2^{e+2}]$ such that $[4]K' = \ker(F)$. To compute F and extract $E_{\mathfrak{a}}$ from its codomain, we take as input:

- The points P_u, Q_u, P_v, Q_v computed in Step 2.
- The integers $N_1, N_2, g_u, x_u, y_u, g_v, x_v, y_v, e$ obtained from a solution of Equation (10) obtained in Step 1.

This data is not only useful to compute T_1, \ldots, T_4 but also gluing and diagonal isogenies located in the beginning of the 2-isogeny chain F that must be handled with care. For this technical reason, one must provide the data above and not

 T_1, \dots, T_4 or kernel points of F directly. We refer to Appendix **B** for a detailed presentation of the computation of F with level 2 theta coordinates (following the approach of [18]) and to Appendix **A** for a cryptographer friendly introduction to higher dimensional isogeny computations with theta coordinates.

Remark 5.3 (Working over \mathbb{F}_p). Using the basis from Equation (12) for our elliptic curves, we can work with rational theta null points all along our dimension 4 isogeny chain. This is one of the reasons to work on the surface: on the floor, the elliptic curve level 2 theta null points are not rational. Our kernel generators have Frobenius-eigenvalue ± 1 , so define \mathbb{F}_p -rational points on the Kummer variety and thus their Theta-coordinates are \mathbb{F}_p -rational (recall that level-2 theta coordinates only give an embedding of the Kummer into projective space, not of the whole variety). This allows us to do the full dimension 4 isogeny computation over \mathbb{F}_p , with the exception of the gluing isogenies. Indeed, as currently implemented the gluing formulas mix points with eigenvalue 1 and points with eigenvalue -1, so we need to temporally work over \mathbb{F}_p^2 . We expect that using more symmetric gluing formulas would allow us to stay over \mathbb{F}_p even for gluing, but we leave that for future work.

6 Implementation results

In this section we describe different implementation choices and their relative efficiency tradeoffs. We give concrete timings, and compare them with the current state of the art of group action based protocols. Find our implementation at

https://github.com/pegasis4d

6.1 Parameter choices

Since our algorithm relies on higher dimensional representations of isogenies, we start with choosing primes of the form $p = c2^f - 1$, ensuring that the full 2^f -torsion is defined over \mathbb{F}_{p^2} . Furthermore, since the probability that Equation (10) admits a solution, depends heavily on how close 2^f is to p, we restrict to the case where c is small.

Among the possible choices for c we retain those that maximize the number of small primes that split in \mathcal{O} , and in particular, primes that are 3 mod 4, since these will greatly increase the probability of finding \mathfrak{B} -good pairs (u, v).

To choose the set \mathfrak{B} , we proceeded as follows: we decided to only include split primes in \mathfrak{B} , which has 3 advantages: first, we can execute the whole algorithm over \mathbb{F}_p , instead of over \mathbb{F}_{p^2} , resulting in a major speed-up. Second, it allows us to use Elkies' algorithm to compute small degree isogenies without explicitly constructing kernel points over extensions of \mathbb{F}_p . Third: such split primes are the only ones that appear as factors of norms of primitive ideals in \mathcal{O} .

The number of primes to include in \mathfrak{B} a priori looks like a trade-off between Steps 1 and 2: including more primes in \mathfrak{B} makes Equation (10) easier to solve (Step 1 becomes faster), but Step 2 will have more (and larger) Elkies isogenies to compute. However, this needs to be combined with the rerandomization technique introduced in Section 5.1. We minimized the overall running time of Steps 1 and 2, and finally derived the parameter sets given in Table 1.

Parameter set	f	c	B
500	503	33	2, 3, 7, 11, 13
1000	1004	15	2, 3, 5, 7, 11
1500	1551	9	2, 3, 5, 11
2000	2026	51	2, 3, 7, 11, 17
4000	4084	63	2, 3, 7, 11, 17, 19

Table 1. Parameter sets used in our implementation. The prime p is of the form $p = c2^{f} - 1$ and \mathfrak{B} is the set of small split primes used.

6.2 Solving the norm equation

The first step (Section 5.1) begins with creating a reduced basis of the starting ideal \mathfrak{a} , followed by a search for small vectors. We are looking for short vectors of (reduced) norm roughly \sqrt{p} , but we allow them to be slightly larger at this point (up to $\log(p)\sqrt{p}$). When we have a list of candidates, we proceed by factoring out all ideals lying above primes in \mathfrak{B} , and then sort by the norms of the remaining ideals.

We then attempt to solve Equation (10) by taking pairs of the shortest vectors that we found. In general, the more pairs, the higher the probability to find a solution. However, the first combinations will be the smallest ones, hence the ones with a highest probability of success. It makes sense then to only try combinations of the smallest vectors. In practice, we observed that taking combinations only involving the 10 shortest vectors, and rerandomizing upon failure, was the most efficient choice for all parameter sets.

Finally, for each pair of (u, v) that we obtain, we have to attempt to write u/g_u and v/g_v as sums of squares. For this, we do trial division on both with all prime factors up to 10^4 for the 500 and 1000 bits parameters, and up to 10^5 for all larger parameters. If we find factors 3 mod 4 (with odd exponent) not included in \mathfrak{B} , we immediately discard the pair. Otherwise, we test what is left for primality. If they are prime, we have the factorization of (u, v) and hence their decomposition as sum of two squares. Otherwise, we move on to the next pair.

6.3 Results and Summary

The results of our SageMath 10.5 implementation can be found in Table 2; timings for each steps are in seconds, and are obtained by averaging 100 runs on an Intel Core i5-1235U clocked at 4.0 GHz.

Parameter set	Step 1	Step 2	Step 3	Tot. Time
500	$0.097~{\rm s}$	$0.48 \mathrm{\ s}$	$0.96 \mathrm{~s}$	$1.53 \mathrm{~s}$
1000	$0.21 \mathrm{~s}$	$1.16 \mathrm{~s}$	$2.84~\mathrm{s}$	$4.21 \mathrm{\ s}$
1500	$1.19 \mathrm{~s}$	$2.85~\mathrm{s}$	$6.49 \mathrm{~s}$	$10.5 \mathrm{\ s}$
2000	$1.68 \mathrm{~s}$	$8.34~\mathrm{s}$	$11.3 \mathrm{~s}$	$21.3~\mathrm{s}$
4000	$15.6~{\rm s}$	$52.8~\mathrm{s}$	$53.5~\mathrm{s}$	$122 \mathrm{~s}$

Table 2. SageMath 10.5 timings on Intel Core i5-1235U at 4.0 GHz, where s denotes the number of seconds in wall-clock time. Step 1 is the time used to solve the norm equation, Step 2 is the time used to derive the kernel of the dimension 4 isogeny, and Step 3 is the time used to compute the dimension 4 isogeny.

We conclude this section with a more detailed comparison with the other available isogeny based EGAs in the literature. The comparison is summarised in Table 3. The timings for SCALLOP [21] were reportedly measured on an Intel i5-6440HQ processor clocked at 3.5 GHz, while the timings for SCALLOP-HD [15] were reportedly measured on an Intel Alder Lake CPU core clocked at 2.1 GHz, and the timings for PEARL-SCALLOP [3] was measured on an Intel i5-1038NG7 processor clocked at 2.0 GHz. The timings for the two versions of KLaPoTi [49] were all re-measured on the same hardware setup as the timings presented in Table 2.

PEGASIS is the first instantiation of an EGA at the 2000 bits and 4000 bits security level. In fact, it is even the first time that the full CSIDH group action can be computed at the 1000 bits level. However, as Table 3 shows, PEGASIS also significantly outperforms all earlier works at their security levels. The closest comparison comes with the Rust implementation of KLaPoTi. However, the fact that KLaPoTi was able to achieve a speedup by two orders of magnitude simply by switching from a high-level SageMath implementation to a low-level Rust implementation is also very promising for PEGASIS. Under the assumption that a comparable speedup would be possible for PEGASIS, we see that PEGASIS gives a highly-practical EGA, even at the highest security levels.

One of the reasons of this efficiency gain is that unlike all the other EGA instantiations, we are able to use the orientation given by Frobenius. This has three key benefits:

- Evaluating the orientation is very efficient,
- We do not need to represent, nor push forward the orientation,
- We can work over \mathbb{F}_p .

Paper	Impl.	500	1000	1500	2000	4000
SCALLOP [21]*	C++	$35 \mathrm{~s}$	$750 \mathrm{~s}$	_	_	_
SCALLOP-HD [15]*	Sage	88 s	$1140~{\rm s}$	_	_	_
PEARL-SCALLOP [3]*	C++	$30 \mathrm{~s}$	$58 \ s$	$710 \mathrm{~s}$	-	_
KL aDoT; [40]	Sage	$207 \mathrm{~s}$	_	_	-	_
KLai 011 [49]	Rust	$1.95~{\rm s}$	-	-	-	_
PEGASIS (This work)	Sage	$1.53 \mathrm{~s}$	$4.21~\mathrm{s}$	$10.5~{\rm s}$	$21.3~{\rm s}$	$121 \mathrm{~s}$

Table 3. Comparison between PEGASIS and other effective group actions in the literature. The last 5 columns gives the timings corresponding to the different security levels, where s gives the number of seconds in wall-clock time. The SCALLOP variants are starred because they were measured on a different hardware setup.

The price to pay is the need to go up to dimension 4 to compute the action, but as Table 2 show that it is still reasonably efficient. We stress that our implementation is only a proof of concept, to explore whether level 4000 was feasible in practice. In particular, we are missing many standard implementation tricks; as a simple example, we work with affine coordinates instead of projective coordinates. Despite this, our timings are very encouraging, and we hope a low level optimised implementation could even be made comparable in efficiency to a state-of-the-art implementation of CSIDH ran as a REGA, like in [12].

6.4 Further improvements

As described in Section 4.2, it is possible to force only one among u/g_u and v/g_v to be a sum of squares, and perform the other isogeny in dimension 4. Note that such 4-dimensional isogeny will have only half of the length of the isogeny that we perform at the end, thus being twice as fast (at least, since the isogeny computation is quasi-linear in its length). On the other hand, Steps 1 and 2 would both become much faster. This approach hence becomes promising as soon as the 4-dimensional isogeny computation has a cost comparable to Step 2, which is the case for the 2000 and 4000 parameter (see Table 2). For such parameters, we computed the first two steps of the algorithm, with $\mathfrak{B} = \{2, 3, 7\}$ in both cases; the results are reported in Table 4. Estimating the cost of one u-isogeny in dimension 4 as half of the cost of Step 3 in Table 2, we expect to reach 21.3 seconds for the 2000 parameter and 106.2 seconds for the 4000. This approach is hence comparable to ours in the former case, and even faster in the latter. We leave its implementation as future work. On the other hand, doing both u and v using 4-dimensional isogenies is less efficient at all security levels we have considered.

Parameter	Step 1	Step 2	Total time	N. Rerand.
2000	0.49 s	$3.83 \mathrm{~s}$	4.32 s	0.70
4000	$3.25 \mathrm{~s}$	$22.8~\mathrm{s}$	$26.0~\mathrm{s}$	1.25

Table 4. Timings for Step 1 and Step 2 when solving the norm equation with single sum of squares, where s again denotes seconds in wall-clock time. Additionally the last column gives the average number of rerandomisations needed.

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A An introduction to efficient higher dimensional isogeny computation

In this section, we explain how higher dimensional isogenies are computed with the theta model. We use the algorithms introduced in [18] for SQIsignHD [19] and the attacks against SIDH. In Appendix A.1, we first introduce some basic theory on theta coordinates in a cryptographer friendly way. We then briefly explain how to change theta coordinates to obtain a system of theta coordinates adapted to a 2^{e} -isogeny, how such an isogeny is computed and how to obtain theta coordinates from Montgomery (x : z)-coordinates on a product of Montgomery elliptic curves. In Appendix A.2, we give more details on changes of theta coordinates adapted to an isogeny obtained from Kani's lemma and how to decompose the codomain into a product.

A.1 Some theory on theta functions

We recall some basics on the theta model. We refer to [20, § 2.1] for a cryptographer friendly introduction. Let (A, λ) be a principally polarised abelian variety of dimension g over an algebraically closed field k. The polarisation $\lambda : A \to \widehat{A}$ is induced by an ample divisor D on A as follows $\lambda = \lambda_D : A \to \widehat{A}, x \mapsto [t_x^*D - D]$. Without loss of generality, we can assume that D is symmetric $[-1]^*D \sim D$. Since λ is a principal polarisation, $\ker(\lambda) = \{0\}$. We define the polarisation of level n as $\lambda \circ [n] = \lambda_{nD} : x \mapsto [t_x^*nD - nD]$ whose kernel is $\ker(\lambda \circ [n]) = A[n]$.

Theta-structures. Formally, a theta-stucture Θ_n of level n on $(A, \lambda \circ [n])$ is an isomorphism between the Heisenberg group $\mathcal{H}(n)$ of level n and the theta-group $\mathcal{G}(nD)$ of nD, $\Theta_n : \mathcal{G}(nD) \xrightarrow{\sim} \mathcal{H}(n)$, that we do not define here. However, this level of abstraction is not needed in the following. The reader only needs to know that a theta structure Θ_n induces a system of coordinates $(\theta_i)_{i \in (\mathbb{Z}/n\mathbb{Z})^g}$ (where $g = \dim(A)$) called theta-coordinates. They define a map into the projective space $A \rightarrow \mathbb{P}_k^{n^g-1}, x \mapsto (\theta_i(x))_{i \in (\mathbb{Z}/n\mathbb{Z})^g}$ which is an embedding when $n \geq 3$ [43, p. 163] and induces an embedding of the Kummer variety $A/\pm \hookrightarrow \mathbb{P}_k^{2^g-1}$ when n = 2 and A is not a product [9, Theorem 4.8.1]. In practice, we work in level n = 2 so theta-coordinates represent points on the Kummer variety *i.e.* points up to a sign.

A polarised abelian variety with a level n theta structure $(A, \lambda \circ [n], \Theta_n)$ is generally too much data to represent on a computer. Usually, we represent it by its associated *theta null point* $(\theta_i(0))_{i \in (\mathbb{Z}/n\mathbb{Z})^g}$. When n = 2 we do not have a guarantee that the theta null point fully determines $(A, \lambda \circ [n], \Theta_n)$. However, this information is enough in practice to perform arithmetic computations on a polarised abelian variety $(A, \lambda \circ [2])$ that is neither a product nor 2-isogenous to a product. These arithmetic operations include doubling $x \mapsto 2x$ and differential addition $x, y, x - y \mapsto x + y$ (see [52, Algorithm 4.4.10]). **Change of theta-coordinates.** A symmetric theta structure of level n is fully determined by a symplectic basis of A[2n] [42, Remark 3, p. 319]. Such a basis $(S_1, \dots, S_q, T_1, \dots, T_q)$ of A[2n] satisfies:

$$\forall 1 \le j \le n, \quad e_{2n}(S_i, S_j) = e_{2n}(T_i, T_j) = 1 \quad \text{and} \quad e_{2n}(T_i, T_j) = \zeta^{\delta_{i,j}},$$

where e_{2n} is the 2*n*-th Weil pairing and $\zeta \in k$ is a primitive 2*n*-th root of unity. Such a basis is called a ζ -symplectic basis. In the following, we shall always assume that theta structures are *symmetric*.

A change of symplectic basis induces a change of theta structure, hence a change of theta coordinates. Namely, let Θ_n and Θ'_n be two theta-structures and respectively induced by ζ -symplectic basis \mathscr{B} and \mathscr{B}' of A[2n]. Let $M \in \operatorname{Sp}_{2g}(\mathbb{Z}/n\mathbb{Z})$ be the change of basis matrix from \mathscr{B} to \mathscr{B}' . Let $(\theta_i)_i$ and $(\theta'_i)_i$ be the systems of theta-coordinates associated to Θ_n and Θ'_n respectively. Then, there is an explicit matrix $N(M, \zeta) \in GL_n(k)$ such that $(\theta'_i)_i = N(M, \zeta) \cdot (\theta_i)_i$ that can be computed as follows.

Theorem 10. [18, Theorem 12] Let us write the symplectic change of basis matrix from \mathscr{B} to \mathscr{B}' by $g \times g$ blocks:

$$M := \begin{pmatrix} A & C \\ B & D \end{pmatrix}$$

Then, there exists $i_0 \in (\mathbb{Z}/n\mathbb{Z})^g$ and $\lambda \in k^*$ such that for all $i \in (\mathbb{Z}/n\mathbb{Z})^g$,

$$\theta_i' = \lambda \sum_{j \in (\mathbb{Z}/n\mathbb{Z})^g} \zeta^{\langle i|j \rangle - \langle Ai + Cj + 2i_0|Bi + Dj \rangle} \theta_{Ai + Cj + i_0},$$

where $\langle i|j \rangle := \sum_{k=1}^{g} i_k j_k$ is the usual scalar product. We can choose any value of $i_0 \in (\mathbb{Z}/n\mathbb{Z})^g$ such that

$$\sum_{\in (\mathbb{Z}/n\mathbb{Z})^g} \zeta^{-\langle Cj+2i_0|Dj\rangle} \theta_{i_0+Cj} \neq 0.$$

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Isogeny computations. Let $F : A \to B$ be a *d*-isogeny. If $\mathscr{B} = (S_1, \dots, S_g, T_1, \dots, T_g)$ is a ζ -symplectic basis of A[4d] adapted to F i.e. such that $\ker(F) = [4]\langle T_1, \dots, T_g \rangle$, then $\mathscr{C} = ([d]F(S_1), \dots, [d]F(S_g), F(T_1), \dots, F(T_g))$ is a ζ^d -symplectic basis of B[4] [19, Theorem 56]. In particular, it means that if we work with theta coordinates $(\theta_i^A)_{i \in (\mathbb{Z}/2\mathbb{Z})^g}$ induced by the symmetric theta structure associated to $[d]\mathscr{B}$ on the domain, then the codomain theta coordinates $(\theta_i^B)_{i \in (\mathbb{Z}/2\mathbb{Z})^g}$ will be induced by the symmetric theta structure associated to \mathscr{C} .

When d = 2, [18, § 4.2] provides formulas to compute the codomain theta null point $(\theta_i^B(0))_{i \in (\mathbb{Z}/2\mathbb{Z})^g}$ from the theta coordinates of T_1, \dots, T_g . The codomain theta null point $(\theta_i^B(0))_i$ is then sufficient to evaluate F *i.e.* to compute $(\theta_i^B(F(x)))_i$ given $(\theta_i^B(x))_i$ for all $x \in A(k)$. Hence the codomain theta null point not only represents the codomain B but also the 2-isogeny F itself. When we say that we compute F, we usually say that we compute its codomain theta null point (and relevant information trivially derived from it). Note the main difference with Vélu's formulas: 8-torsion points T_1, \dots, T_g are necessary for this computation; 4 or 2-torsion points would not be sufficient.

When $d = 2^e$, and $F = f_e \circ \cdots \circ f_1$ is a chain of 2-isogenies, a choice of symplectic basis \mathscr{B} adapted to F (*i.e.* such that ker $(F) = [4]\langle T_1, \cdots, T_g \rangle$) induces a choice of level 2 theta-structure on the codomain of f_i associated to $[2^{e-i}]f_i \circ \cdots \circ f_1([2^i]S_1, \cdots, [2^i]S_g, T_1, \cdots, T_g)$ for all $i \in [1; e]$, so that we can propagate the formulas from $[18, \S 4.2]$ along the chain.

When F has been obtained via Kani's lemma, it is defined on a product of abelian varieties (e.g. a product of elliptic curves) so the first steps of the chain are gluing isogenies $f : A_1 \times A_2 \to B$ that need to be handled with extra care since the formulas are different to compute them. Splitting isogenies along the chain $f : A \to B_1 \times B_2$ also require some attention because point duplications $x \mapsto 2x$ are more difficult on their domain and codomain than generic ones. In particular, we need to detect these gluings and splittings in advance. This will be the main focus of Appendix B in our special case of interest (when F is the isogeny from Section 5.3).

As in dimension 1, divide and conquer strategies can be applied to minimize the number of duplications and isogeny evaluations required to compute a 2isogeny chain $F = f_e \circ \cdots \circ f_1$. These strategies can also be adapted to take into account the specific constraints imposed by gluing and splitting isogenies and their relative cost. With 4-dimensional computations in mind, more details on these strategies can be found in [18, Appendix E].

Product theta-structures. When we compute isogenies obtained via Kani's lemma, we start and terminate on a product of abelian varieties which are naturally equiped with a *product theta-structure*. If $A = A_1 \times A_2$ and $(A_1, \lambda_1), (A_2, \lambda_2)$ are principally polarised abelian varieties, we consider the product polarisation $\lambda := \lambda_1 \times \lambda_2 : (x, y) \mapsto (\lambda_1(x), \lambda_2(x))$. If $\Theta_n^{A_i}$ is a level *n* theta-structure on $(A_i, \lambda_i \circ [n])$ induced by a ζ -symplectic basis $\mathscr{B}_i = (S_1^{(i)}, \cdots, S_{g_i}^{(i)}, T_1^{(i)}, \cdots, T_{g_i}^{(i)})$ of A[2n] and of associated system of theta-coordinates $(\theta_j^{A_i})_{j \in (\mathbb{Z}/n\mathbb{Z})^{g_i}}$ for $i \in \{1, 2\}$, then we can define the product theta-structure $\Theta_n := \Theta_n^{A_1} \times \Theta_n^{A_2}$ on (A, λ) as follows. It is the level *n* theta-structure induced by the ζ -symplectic basis of A[2n]:

$$\mathscr{B} = \mathscr{B}_1 \times \mathscr{B}_2 := ((S_1^{(1)}, 0), \cdots, (S_{g_1}^{(1)}, 0), (0, S_1^{(2)}), \cdots, (0, S_{g_2}^{(2)}), (T_1^{(1)}, 0), \cdots, (T_{g_1}^{(1)}, 0), (0, T_1^{(2)}), \cdots, (0, T_{g_2}^{(2)})).$$
(15)

The theta-coordinates $(\theta_j^A)_{j \in (\mathbb{Z}/n\mathbb{Z})^{g_1+g_2}}$ associated to $\Theta_n = \Theta_n^{A_1} \times \Theta_n^{A_2}$ are given as follows:

$$\forall (x,y) \in A_1(k) \times A_2(k), (j_1, j_2) \in (\mathbb{Z}/n\mathbb{Z})^{g_1+g_2}, \quad \theta^A_{j_1, j_2}(x,y) = \theta^{A_1}_{j_1}(x)\theta^{A_2}_{j_2}(y).$$
(16)

In the context of a 2^e-isogeny computation $F : A \to B$ $(n = 2^{e+2})$, we usually start with a product theta-structure induced by \mathscr{B} and we have to compute a change of theta-coordinates induced by a change of symplectic basis from \mathscr{B} to $\mathscr{B}' = (S_1, \cdots, S_q, T_1, \cdots, T_q)$ such that ker $(F) = [4]\langle T_1, \cdots, T_q \rangle$.

The codomain is also often a product $B = B_1 \times B_2$ but the level 2 theta-structure on the codomain induced by F is associated to $\mathscr{C} = ([2^e]F(S_1), \cdots, [2^e]F(S_g), F(T_1), \cdots, F(T_g))$ which is not a product theta-structure. A change of theta-coordinates is necessary to recover the product theta-coordinates on B. Once this has been done, we obtain the *theta-null point* of the product $(\theta_{j_1,j_2}^B(0))_{j_1,j_2} = (\theta_{j_1}^{B_1}(0) \cdot \theta_{j_2}^{B_2}(0))_{j_1,j_2}$ from which we can infer the theta-null points $(\theta_j^{B_i}(0))_j$ of each component B_i , which can then be used to determine the polarised abelian variety $(B_i, \lambda_{B_i} \circ [2])$. In Appendix A.2, we shall see how to compute the necessary change of theta-coordinates for this splitting operation when F is obtained from Kani's lemma and apply this method to our case of interest (when F is the isogeny from Section 5.3) in Appendix B.4.

Theta-structures on Montgomery curves. In our context, we work with isogenies between products of elliptic curves. These elliptic curves are equipped with a Montgomery model over \mathbb{F}_{p^2} (and even \mathbb{F}_p for CSIDH) so we need to convert this model into a level 2 theta model. Conversely, we also need to translate a level 2 theta model obtained after the splitting operation mentioned above into a Montgomery model. The focus of this paragraph will be to explain these conversions.

If E is a Montgomery curve, then Montgomery (x : z)-coordinates and level 2 theta coordinates $(\theta_0 : \theta_1)$ both induce *Kummer lines i.e.* isomorphisms $E/\pm \Rightarrow \mathbb{P}^1$. Hence, a change of coordinates between theta and Montgomery is a homography. In [54, Chapter 7, Appendix A], formulas were introduced to express such a homography when the theta structure is symmetric and associated to a 4-torsion basis of specific shape.

Let *E* be an elliptic curve in the Montgomery model and (P,Q) be a basis of E[4] such that Q := (-1:1) (in (x:z) coordinates). Let us write P := (r:s). Then, we may define a level 2 theta-structure on *E* with theta-null point (a:b) := (r+s:r-s). The conversion map from Montgomery to theta coordinates is then $(x:z) \mapsto (a(x-z):b(x+z))$.

Conversely, if (a:b) is the theta-null point, the conversion map from theta to Montgomery coordinates is $(\theta_0, \theta_1) \mapsto (a\theta_1 + b\theta_0 : a\theta_1 - b\theta_0)$. Via this map, the 2-torsion theta points (b:a), (a:-b) and (b:-a) are mapped to the 2-torsion Montgomery points $(a^2+b^2:a^2-b^2)$, (0:1) and $(a^2-b^2:a^2+b^2)$ respectively. Identifying the 2-torsion, we can then obtain the Montgomery equation of E:

$$By^{2} = x(x - \alpha)(x - 1/\alpha) = x^{3} + Ax^{2} + x,$$

with $\alpha := (a^2 + b^2)/(a^2 - b^2)$ and $A := -(\alpha + 1/\alpha) = -2(a^4 + b^4)/(a^4 - b^4)$. To identify B, we need to know a full point of 4-torsion (x : y : z) on E. This will be important in our context to distinguish the result of the CSIDH class group action $E_{\mathfrak{a}}$ from its quadratic twist over \mathbb{F}_p (see Appendix B.4).

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As we have seen previously, a fixed Montgomery curve E has several level 2 theta structures related by symplectic matrices between 4-torsion basis. Using Theorem 10, we can then obtain theta structures which are not induced by a 4-torsion basis (P,Q) with $(x(Q) : z(Q)) \neq (-1:1)$. Conversely, from different theta structures on E, we obtain 6 different possible Montgomery coefficients Afrom the conversion formula $(\theta_0, \theta_1) \mapsto (a\theta_1 + b\theta_0 : a\theta_1 - b\theta_0)$.

Proposition 11. Let (a : b) be a level 2 theta null point on a Montgomery curve E associated to a symmetric theta structure. Then all possible Montgomery coefficients of E are of the form:

$$-2\frac{a^4+b^4}{a^4-b^4}, \quad -\frac{a^4+6a^2b^2+b^4}{2(a^3b+b^3a)}, \quad \zeta_4\frac{a^4-6a^2b^2+b^4}{2(a^3b-b^3a)}, \\ 2\frac{a^4+b^4}{a^4-b^4}, \quad \frac{a^4+6a^2b^2+b^4}{2(a^3b+b^3a)}, \quad -\zeta_4\frac{a^4-6a^2b^2+b^4}{2(a^3b-b^3a)}.$$

where ζ_4 is a root of -1. They correspond respectively to theta null points:

$$(a:b), (a+b:a-b), (a+\zeta_4b:a-\zeta_4b), (b:a), (a-b:a+b), (a-\zeta_4b:a+\zeta_4b).$$

Proof. Let (P,Q) be a basis of E[4] inducing a symmetric theta structure of theta null point (a:b) and let $\zeta_4 := e_4(P,Q)$. Then, enumerating all symplectic matrices of $\operatorname{Sp}_2(\mathbb{Z}/4\mathbb{Z})$, we obtain all ζ_4 -symplectic basis (P',Q') of E[4] inducing symmetric level 2 theta structures of theta null points (a':b') obtained from (a:b) by Theorem 10. We can then compute all corresponding Montgomery coefficients $A' := -2(a'^4 + b'^4)/(a'^4 - b'^4)$.

A.2 Theta structures on the domain and codomain of an isogeny obtained from Kani's lemma

In the following lemma, we give an explicit description of a symplectic basis adapted to an isogeny obtained from Kani's lemma (in point (i)). By Theorem 10, this can be used to compute a change of theta coordinates from product theta coordinates on the domain before starting the isogeny computation. We also describe a symplectic basis associated to a product level 2 theta structure on the codomain (in point (ii)) and its relation with the symplectic basis naturally induced by F via [19, Theorem 56] (in point (iii)). Using Theorem 10, the resulting symplectic change of basis matrix can be used to compute the product theta null point and product theta coordinates on the codomain. In particular, we can "extract" components of the codomain.

Lemma 12. Let a and b be odd and coprime integers and d := a+b not divisible by char(k). Consider an (a, b)-isogeny diamond (as defined in Lemma 4) between

PPAVs of dimension g:



and the associated Kani d-isogeny:

$$F := \begin{pmatrix} \varphi & \widetilde{\psi'} \\ -\psi & \widetilde{\varphi'} \end{pmatrix} : A \times B' \to B \times A'.$$

Let $(x_1, \dots, x_g, y_1, \dots, y_g)$ be a ζ -symplectic basis of B[4d]. Let α and β be modular inverses of a and b modulo 4d respectively.

(i) For all $i \in [1; g]$, we denote:

$$S_i := ([-\alpha]\widetilde{\varphi}(y_i), 0), \quad S_{i+g} := (0, [\beta]\psi'(x_i)),$$
$$T_i := (\widetilde{\varphi}(x_i), \psi'(x_i)), \quad T_{i+g} := ([1 - \alpha d]\widetilde{\varphi}(y_i), \psi'(y_i))$$

Then $(S_1, \dots, S_{2g}, T_1, \dots, T_{2g})$ is a ζ -symplectic basis of $(A \times B')[4d]$ adapted to F i.e. such that ker $(F) = [4]\langle T_1, \dots, T_{2g} \rangle$.

(ii) For all $i \in [1; g]$, we denote:

$$U_i := ([d]x_i, 0), \quad U_{i+g} := (0, [d]\psi \circ \widetilde{\varphi}(x_i)),$$
$$V_i := ([d]y_i, 0), \quad V_{i+g} := (0, [d\alpha\beta]\psi \circ \widetilde{\varphi}(y_i)).$$

Then $(U_1, \dots, U_{2g}, V_1, \dots, V_{2g})$ is a product ζ^d -symplectic basis of $(B \times A')[4]$.

(iii) The ζ^d -symplectic basis $([d]F(S_1), \cdots, [d]F(S_{2g}), F(T_1), \cdots, F(T_{2g}))$ of $(B \times A')[4]$ naturally induced by F via [19, Theorem 56] is related to $(U_1, \cdots, U_{2g}, V_1, \cdots, V_{2g})$ by the following formulas. For all $i \in [\![1; g]\!]$:

$$[d]F(S_i) = -V_i + [b]V_{i+g}, \quad [d]F(S_{i+g}) = U_i + [\beta]U_{i+g},$$
$$F(T_i) = U_i, \quad F(T_{i+g}) = [b]V_{i+g},$$

so that:

$$U_{i} = F(T_{i}), \quad U_{i+g} = [bd]F(S_{i+g}) - [b]F(T_{i}),$$
$$V_{i} = F(T_{i+g}) - [d]F(S_{i}), \quad V_{i+g} = [\beta]F(T_{i+g}).$$

Proof. (i) For all principally polarised abelian varieties C and D, we denote λ_C the principal polarisation on C and $\lambda_C \times \lambda_D$ the product (principal) polarisation on $C \times D$. Then, since $(x_1, \dots, x_g, y_1, \dots, y_g)$ is a ζ -symplectic basis of A[4d], we have the following results for the 4*d*-th Weil-pairing associated to the product polarisation $\lambda_A \times \lambda_{B'}$:

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$$\begin{split} e_{4d}^{\lambda_A \times \lambda_{B'}}(S_i, S_j) &= e_{4d}^{\lambda_A} ([-\alpha] \widetilde{\varphi}(y_i), [-\alpha] \widetilde{\varphi}(y_j)) = e_{4d}^{\lambda_A}(y_i, y_j)^{a\alpha^2} = 1 \\ e_{4d}^{\lambda_A \times \lambda_{B'}}(S_i, S_{j+g}) &= e_{4d}^{\lambda_A} ([-\alpha] \widetilde{\varphi}(y_i), 0) e_{4d}^{\lambda_{B'}}(0, [\beta] \psi'(x_j)) = 1 \\ e_{4d}^{\lambda_A \times \lambda_{B'}}(T_i, T_j) &= e_{4d}^{\lambda_A} (\widetilde{\varphi}(x_i), \widetilde{\varphi}(x_j)) e_{4d}^{\lambda_{B'}}(\psi'(x_i), \psi'(x_j)) = e_{4d}^{\lambda_A}(x_i, x_j)^{a+b} = 1 \\ e_{4d}^{\lambda_A \times \lambda_{B'}}(T_i, T_{j+g}) &= e_{4d}^{\lambda_A} (\widetilde{\varphi}(x_i), [1 - \alpha d] \widetilde{\varphi}(y_j)) e_{4d}^{\lambda_{B'}}(\psi'(x_i), \psi'(y_j)) \\ &= e_{4d}^{\lambda_A}(x_i, y_j)^{a(1 - \alpha d)+b} = e_{4d}^{\lambda_A}(x_i, y_j)^0 = 1 \\ e_{4d}^{\lambda_A \times \lambda_{B'}}(S_i, T_j) &= e_{4d}^{\lambda_A} ([-\alpha] \widetilde{\varphi}(y_i), \widetilde{\varphi}(x_j)) e_{4d}^{\lambda_{B'}}(0, \psi'(x_j)) = e_{4d}^{\lambda_A}(x_j, y_i)^{a\alpha} = \zeta^{\delta_{i,j}} \\ e_{4d}^{\lambda_A \times \lambda_{B'}}(S_i, T_j) &= e_{4d}^{\lambda_A} ([-\alpha] \widetilde{\varphi}(y_i), [1 - \alpha d] \widetilde{\varphi}(y_j)) e_{4d}^{\lambda_{B'}}(0, \psi'(y_j)) \\ &= e_{4d}^{\lambda_A}(y_i, y_j)^{-(1 - \alpha d)\alpha} = 1 \\ e_{4d}^{\lambda_A \times \lambda_{B'}}(S_{i+g}, T_j) &= e_{4d}^{\lambda_A}(0, \widetilde{\varphi}(x_j)) e_{4d}^{\lambda_{B'}} ([\beta] \psi'(x_i), \psi'(x_j)) = e_{4d}^{\lambda_A}(x_i, x_j)^{b\beta} = 1 \\ e_{4d}^{\lambda_A \times \lambda_{B'}}(S_{i+g}, T_j) &= e_{4d}^{\lambda_A}(0, [1 - \alpha d] \widetilde{\varphi}(y_j)) e_{4d}^{\lambda_{B'}}([\beta] \psi'(x_i), \psi'(y_j)) \\ &= e_{4d}^{\lambda_A}(x_i, y_j)^{b\beta} = e_{4d}^{\lambda_A}(x_i, y_j) = \zeta^{\delta_{i,j}}, \\ \text{for all } i, j \in [1 ; g]. \text{ Hence, } (S_1, \cdots, S_{2g}, T_1, \cdots, T_{2g}) \text{ is a } \zeta\text{-symplectic basis of } (A \times B') [4d]. \text{ Since } ([4]x_1, \cdots, [4]x_g, [4]x_1, \cdots, [4]y_g) \text{ generates } A[d], \text{ we clearly have by Lemma 4:} \\ [4] \langle T_1, \cdots, T_{2g} \rangle = \{([a]x, \psi' \circ \varphi(x)) \mid x \in A[d]\} = \ker(F). \\ \text{This proves (i)} \end{split}$$

This proves (i). (ii) $\mathscr{B} := (U_1, \cdots, U_{2g}, V_1, \cdots, V_{2g})$ is the product $\mathscr{B}_1 \times \mathscr{B}_2$ with

$$\mathscr{B}_1 := ([d]x_1, \cdots, [d]x_g, [d]y_1, \cdots, [d]y_g)$$

 $\mathscr{B}_{2} := ([d]\psi \circ \widetilde{\varphi}(x_{1}), \cdots, [d]\psi \circ \widetilde{\varphi}(x_{g}), [d\alpha\beta]\psi \circ \widetilde{\varphi}(y_{1}), \cdots, [d\alpha\beta]\psi \circ \widetilde{\varphi}(y_{g})).$

And we have for all $i,j\in [\![1\ ;\ g]\!]:$

$$e_4^{\lambda_B}([d]x_i, [d]x_j) = e_{4d}^{\lambda_B}(x_i, x_j)^d = 1$$

$$e_4^{\lambda_B}([d]y_i, [d]y_j) = e_{4d}^{\lambda_B}(y_i, y_j)^d = 1$$
$$e_4^{\lambda_B}([d]x_i, [d]y_j) = e_{4d}^{\lambda_B}(x_i, y_j)^d = \zeta^{d\delta_{i,j}}$$

so \mathscr{B}_1 is a ζ^d -symplectic basis of B[4]. Similarly, we verify that \mathscr{B}_2 is a ζ^d -symplectic basis of A'[4]. It follows that $\mathscr{B} = \mathscr{B}_1 \times \mathscr{B}_2$ is a product ζ^d -symplectic basis of $(B \times A')[4]$, as desired.

(iii) The fact that $([d]F(S_1), \dots, [d]F(S_{2g}), F(T_1), \dots, F(T_{2g}))$ is a ζ^d -symplectic basis of $(B \times A')[4]$ follows from [19, Theorem 56]. It remains to compute for all $i \in [1; g]$:

$$\begin{split} [d]F(S_i) &= [d]F([-\alpha]\widetilde{\varphi}(y_i), 0) = ([-d\alpha]\varphi \circ \widetilde{\varphi}(y_i), [d\alpha]\psi \circ \widetilde{\varphi}(y_i)) \\ &= ([-d\alpha\alpha]y_i, [d\alpha]\psi \circ \widetilde{\varphi}(y_i)) = -V_i + [b]V_{i+g} \end{split}$$

$$[d]F(S_{i+g}) = [d]F((0, [\beta]\psi'(x_i)) = ([d\beta]\widetilde{\psi}' \circ \psi'(x_i), [d\beta]\widetilde{\varphi}' \circ \psi'(x_i))$$

$$= ([d\beta b]x_i, [d\beta]\psi \circ \widetilde{\varphi}(x_i)) = ([d]x_i, [d\beta]\psi \circ \widetilde{\varphi}(x_i)) = U_i + [\beta]U_{i+g}$$

$$\begin{split} F(T_i) &= F(\widetilde{\varphi}(x_i), \psi'(x_i)) = (\varphi \circ \widetilde{\varphi}(x_i) + \widetilde{\psi'} \circ \psi'(x_i), -\psi \circ \widetilde{\varphi}(x_i) + \widetilde{\varphi'} \circ \psi'(x_i)) \\ &= ([a+b]x_i, -\psi \circ \widetilde{\varphi}(x_i) + \psi \circ \widetilde{\varphi}(x_i)) = ([d]x_i, 0) = U_i \end{split}$$

$$\begin{split} F(T_{i+g}) &= F([1 - \alpha d]\widetilde{\varphi}(y_i), \psi'(y_i)) \\ &= ([1 - \alpha d]\varphi \circ \widetilde{\varphi}(y_i) + \widetilde{\psi'} \circ \psi'(y_i), -[1 - \alpha d]\psi \circ \widetilde{\varphi}(y_i) + \widetilde{\varphi'} \circ \psi'(y_i)) \\ &= ([a(1 - \alpha d) + b]y_i, [\alpha d - 1 + 1]\psi \circ \widetilde{\varphi}(y_i)) = (0, [d]\psi \circ \widetilde{\varphi}(y_i)) = [b]V_{i+g}, \end{split}$$

where we used twice the fact that $\psi' \circ \varphi = \varphi' \circ \psi$ implies that $\widetilde{\varphi'} \circ \psi' = \psi \circ \widetilde{\varphi}$. The inverse equations:

$$U_{i} = F(T_{i}), \quad U_{i+g} = [bd]F(S_{i+g}) - [b]F(T_{i}),$$
$$V_{i} = F(T_{i+g}) - [d]F(S_{i}), \quad V_{i+g} = [\beta]F(T_{i+g}),$$

follow immediately. This completes the proof.

B Implementing the 4-dimensional isogeny computation for the CSIDH group action in the simplest case

In this Section, we explain how to compute the 4-dimensional isogeny $F: E_u^2 \times E_v^2 \to E_a^2 \times E'^2$ from Section 5.3. We compute F as a chain of e 2-isogenies, using the algorithms from [18, § 4]. The general approach is very similar to SQIsignHD verification and SIDH attacks described in [18, § 5 and Appendix B].

B.1 Where are the gluings and splittings?

Because the formulas for gluing isogenies are different, we need to handle gluings with special care and locate them in advance in the 2-isogeny chain. We have the following result, very similar to [18, Lemma 23]:

Lemma 13. Without loss of generality, we can assume that $2|y_u|$ and $2|y_v|$. Let $m := v_2(x_vy_u - x_uy_v)$. Then F is of the form $F = G \circ F_{m+1} \circ F_m \circ \cdots \circ F_1$, where

$$E_u^2 \times E_v^2 \xrightarrow{F_1} A_1^2 \xrightarrow{F_2} \cdots \xrightarrow{F_{m-1}} A_{m-1}^2 \xrightarrow{F_m} A_m^2 \xrightarrow{F_{m+1}} B_m$$

is a chain of 2-isogenies, the A_i are principally polarised abelian surfaces and Bis a principally polarised abelian variety of dimension 4. For all, $i \in [\![2; m]\!]$, F_i is a diagonal isogeny $\text{Diag}(f_i, f_i)$ with $f_i : A_{i-1} \to A_i$, a 2-dimensional 2-isogeny and $F_1 : (R_1, S_1, R_2, S_2) \mapsto (f_1(R_1, R_2), f_1(S_1, S_2))$ with $f_1 : E_u \times E_v \to A_1$, a gluing 2-isogeny.

Besides, the 2-dimensional 2^m -isogeny $\Phi := f_m \circ \cdots \circ f_1$ has kernel:

 $\ker(\varPhi) = \{([N_1 x_u]\varphi_u(P), [g_u(x_u x_v + y_u y_v)]\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(P)) \mid P \in E_1[2^m]\}.$

Proof. By Eq. (14), we have:

$$\begin{split} \ker(F) &= \{ ([uN_1]\varphi_u(P), [uN_1]\varphi_u(Q), \\ & [g_u]M_v \cdot M_u^T(\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(P), \varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(Q))) \mid P, Q \in E_u[2^e] \}, \end{split}$$

so the first m isogenies of the 2-isogeny chain have kernel:

$$\begin{split} \ker(F_m \circ \dots \circ F_1) &= [2^{e-m}] \ker(F) \\ &= \{([uN_1]\varphi_u(P), [uN_1]\varphi_u(Q), \\ & [g_u(x_ux_v + y_uy_v)]\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(P) \\ &+ [g_u(x_vy_u - x_uy_v)]\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(Q), \\ &- [g_u(x_vy_u - x_uy_v)]\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(P) \\ &+ [g_u(x_ux_v + y_uy_v)]\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(Q)) \mid P, Q \in E_1[2^m] \} \\ &= S(K \times K), \end{split}$$

with $S: E_u^2 \times E_v^2 \to (E_u \times E_v)^2$, $(R_1, S_1, R_2, S_2) \mapsto (R_1, R_2, S_1, S_2)$ and

$$K := \{([N_1 x_u]\varphi_u(P), [g_u(x_u x_v + y_u y_v)]\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k}(P)) \mid P \in E_1[2^m]\}.$$

It follows that $(F_m \circ \cdots \circ F_1) \circ S^{-1}$ is a diagonal 2^m -isogeny

$$\operatorname{Diag}(\varPhi,\varPhi):(E_u\times E_v)^2\to A_m^2$$

with $\ker(\Phi) = K$, so the F_i and F have the desired form.

Locating splittings in the isogeny chain computation is also important. Indeed, on abelian varieties 2-isogenous to a product of abelian varieties, the generic doubling procedure [18, Algorithm 8] may fail because of zero divisions. Doublings are involved to obtain kernel information at every step of the isogeny chain but we have some flexibility on the abelian variety where these doublings take place along the chain, depending on the computational strategy we use. As in [18, Appendix E], we use computational strategies that do not involve doublings on codomains of splitting isogenies. Using the structure of the polarised dual \tilde{F} , we obtain the

Lemma 14. Assume that $2|y_u, y_v|$ and let $m := v_2(x_vy_u - x_uy_v)$ as in Lemma 13. Then F is of the form $F = F_e \circ \cdots \circ F_{e-m} \circ G'$, where

$$B' \xrightarrow{F_{e-m}} A_{e-m}^2 \xrightarrow{F_{e-m+1}} A_{e-m+1}^2 \xrightarrow{F_{e-m+2}} \cdots \xrightarrow{F_{e-m+2}} A_{e-1}^2 \xrightarrow{F_e} E_{\mathfrak{a}}^2 \times (E')^2$$

is a chain of 2-isogenies, the A_i are principally polarised abelian surfaces and B' is a principally polarised abelian variety of dimension 4. For all, $i \in [e-m+1; e]$, F_i is a diagonal isogeny $\operatorname{Diag}(f_i, f_i)$ with $f_i : A_{i-1} \to A_i$, a 2-dimensional 2-isogeny and $F_e := S' \circ \operatorname{Diag}(f_e, f_e)$ with $f_1 : A_{e-1} \to E_{\mathfrak{a}} \times E'$, a splitting 2-isogeny and $S' : (E_{\mathfrak{a}} \times E')^2 \to E_{\mathfrak{a}}^2 \times E'^2$, $(R_1, R_2, S_1, S_2) \mapsto (R_1, S_1, R_2, S_2)$.

Proof. The polarised dual $\widetilde{F}: E^2_{\mathfrak{a}} \times (E')^2 \to E^2_u \times E^2_v$ is given by:

$$\widetilde{F} = \begin{pmatrix} \Phi_u \circ \widetilde{\Phi}_1 & -\Phi'_v \circ \widetilde{\Phi}'_2 \\ -\Phi_v \circ \widetilde{\Phi}_2 & \Phi'_u \circ \widetilde{\Phi}'_1 \end{pmatrix}$$

and its kernel is:

$$\begin{aligned} \ker(\tilde{F}) &= \{ ([uN_1]P, [uN_1]Q, -\varPhi'_2 \circ \tilde{\varPhi'}_v \circ \varPhi_u \circ \tilde{\varPhi}_1(P, Q)) \mid P, Q \in E_{\mathfrak{a}}[2^e] \} \\ &= \{ ([uN_1]P, [uN_1]Q, -M_v^T \cdot M_u(\varphi'_2 \circ \hat{\varphi}'_v \circ \varphi_u \circ \hat{\varphi}_1(P), \\ \varphi'_2 \circ \hat{\varphi}'_v \circ \varphi_u \circ \hat{\varphi}_1(Q))) \mid P, Q \in E_{\mathfrak{a}}[2^e] \} \\ &= \{ ([uN_1]\varphi_u(P), [uN_1]\varphi_u(Q), \\ &- [x_u x_v + y_u y_v]\varphi'_2 \circ \hat{\varphi}'_v \circ \varphi_u \circ \hat{\varphi}_1(P) \\ &- [x_u y_v - x_v y_u]\varphi'_2 \circ \hat{\varphi}'_v \circ \varphi_u \circ \hat{\varphi}_1(Q), \\ [x_u y_v - x_v y_u]\varphi'_2 \circ \hat{\varphi}'_v \circ \varphi_u \circ \hat{\varphi}_1(Q)) \\ &- [x_u x_v + y_u y_v]\varphi'_2 \circ \hat{\varphi}'_v \circ \varphi_u \circ \hat{\varphi}_1(P) \\ &- [x_u x_v + y_u y_v]\varphi'_2 \circ \hat{\varphi}'_v \circ \varphi_u \circ \hat{\varphi}_1(Q)) \mid P, Q \in E_{\mathfrak{a}}[2^e] \} \end{aligned}$$

It follows that

$$\ker(\widetilde{F})[2^m] = S'(K' \times K'),$$

with:

$$K' = \{([uN_1]P, -[x_ux_v + y_uy_v]\varphi_2' \circ \widehat{\varphi}_v' \circ \varphi_u \circ \widehat{\varphi}_1(P)) \mid P \in E_{\mathfrak{a}}[2^m]\}$$

We then easily conclude that F has the desired form.

By Lemma 13, to compute F, we first have to compute a 2-dimensional 2isogeny chain $\Phi := f_m \circ \cdots \circ f_1 : E_u \times E_v \to A_m$, then a 4-dimensional gluing of abelian surfaces $F_{m+1} : A_m^2 \to B$ and e-m-1 generic 4-dimensional 2-isogenies. Except for the first m steps and splitting at the end, we do not have a way of detecting products of abelian varieties located elsewhere in the 2-isogeny chain. However, they can appear with very low probability $\tilde{O}(1/p)$ so we can assume there are none and treat the e-m-1 last 2-isogenies as generic ones.

Hence, we proceed as follows to compute F:

- 1. We compute the 2-dimensional isogeny $\Phi := f_m \circ \cdots \circ f_1 : E_u \times E_v \to A_m$ (Appendix B.2).
- 2. We compute the gluing isogeny $F_{m+1}: A_m^2 \to B$ (Appendix B.3).
- 3. We compute the last e m 1 2-isogenies of the chain as generic isogenies with a computational strategy avoiding doublings m steps before the end, as in [18, Algorithm 21].
- 4. We compute a change of theta-coordinates on the codomain $E_{\mathfrak{a}}^2 \times E'^2$ to obtain the product theta-structure and we retrieve $E_{\mathfrak{a}}$ (Appendix B.4).

In the following, we assume we are given the following inputs defined in Section 5.3:

- The points $P_u, Q_u, P_v, Q_v;$
- The integers $N_1, N_2, g_u, x_u, y_u, g_v, x_v, y_v, e$ obtained from a solution of Eq. (10);

where (P_u, Q_u) and (P_v, Q_v) are images of a basis (P, Q) of $E_1[2^{e+2}]$ via the g_u -isogeny $\varphi_u : E_1 \to E_u$ and the $g_v N_1 N_2$ -isogeny $\varphi_v \circ \widehat{\varphi}_{\mathfrak{c}_k} \circ \varphi_{\mathfrak{b}_k} : E_1 \to E_v$. In the following, we denote $\zeta := e_{2^{e+2}}(P, Q)$, so that $e_{2^{e+2}}(P_u, Q_u) = \zeta^{g_u}$ and $e_{2^{e+2}}(P_v, Q_v) = \zeta^{g_v N_1 N_2}$.

B.2 The 2-dimensional part

When computing the gluing isogeny $f_1 : E_u \times E_v \to A_1$, we first convert the Montgomery (x : z)-coordinates on $E_u \times E_v$ into product theta coordinates. Let (R_u, S_u) and (R_v, S_v) be basis of $E_u[4]$ and $E_v[4]$ respectively such that $x(S_u) = x(S_v) = -1$ and $e_4(R_u, S_u) = e_4(R_v, S_v) = \zeta^{2^e}$. By Appendix A.1, we can then convert Montgomery (x : z)-coordinates on E_u and E_v into level 2 theta coordinates associated to symmetric theta structures induced by (R_u, S_u) and (R_v, S_v) respectively. Then, the resulting product level 2 theta structure on $E_u \times E_v$ is induced by the ζ^{2^e} -symplectic basis of $(E_u \times E_v)[4]$:

$$\mathscr{B}_0 := ((R_u, 0), (0, R_v), (S_u, 0), (0, S_v)).$$

However, \mathscr{B}_0 is not adapted to f_1 in the sense of Appendix A.1, so we need to find a (non-product) symplectic basis adapted to f_1 and to compute the associated change of theta coordinates with Theorem 10. We can then compute the gluing isogeny f_1 in level 2 theta coordinates using [20, Algorithm 8] and generic isogenies f_2, \dots, f_m using [20, Algorithm 7].

Consider the intermediate ζ^{2^e} -symplectic basis \mathscr{B}_1 of $(E_u \times E_v)[4]$, where:

$$\mathscr{B}_1 := (([2^e]P_u, 0), (0, [2^e]P_v), ([2^e\gamma_u]Q_u, 0), (0, [2^e\gamma_v\eta_1\eta_2]Q_v)),$$

and $\gamma_u, \gamma_v, \eta_1, \eta_2$ are inverses of g_u, g_v, N_1, N_2 modulo 2^{e+2} . Given the expression of the kernel of $\Phi := f_m \circ \cdots \circ f_1$ from Lemma 13, we can easily express the symplectic change of basis matrix M_1 from \mathscr{B}_1 to $[2^m]\mathscr{B}_2$, where \mathscr{B}_2 is a $\zeta^{2^{e-m}}$ symplectic basis of $(E_u \times E_v)[2^{m+2}]$ adapted to Φ (given by Lemma 15). This matrix M_1 is described in Lemma 16. We can also express the change of basis from \mathscr{B}_0 to \mathscr{B}_1 :

$$\begin{pmatrix} a_u & 0 & c_u & 0 \\ 0 & a_v & 0 & c_v \\ b_u & 0 & d_u & 0 \\ 0 & b_v & 0 & d_v \end{pmatrix}$$

where $[2^e]P_u = [a_u]R_u + [b_u]S_u$, $[2^e\gamma_u]Q_u = [c_u]R_u + [d_u]S_u$, $[2^e]P_v = [a_v]R_v + [b_v]S_v$ and $[2^e\gamma_v]Q_v = [c_v]R_v + [d_v]S_v$. The change of basis from \mathscr{B}_0 to $[2^m]\mathscr{B}_2$ is then $M_2 := M_0 \cdot M_1$ to which we can directly apply Theorem 10 to obtain the desired change of theta coordinates.

Lemma 15. We keep the notations from Appendix B.1 and Lemma 13. Let $\zeta := e_{2^{e+1}}(P,Q), s_{uv} := x_u x_v + y_u y_v, \delta_{uv} := x_v y_u - x_u y_v, \eta_1, \eta_2, \gamma_u, \gamma_v, \mu, \sigma_{uv}$ be inverses of $N_1, N_2, g_u, g_v, u = g_u(x_u^2 + y_u^2), s_{uv}$ modulo 2^{m+2} respectively. Then, $\mathscr{B}_2 := (J_1, J_2, K_1, K_2)$ given by:

$$J_{1} := \left(\left[-2^{e-m} \eta_{1} \mu \gamma_{u} \right] Q_{u}, 0 \right), \quad J_{2} := \left(0, \left[2^{e-m} \gamma_{u} \gamma_{v} \eta_{1} \eta_{2} \sigma_{uv} \right] P_{v} \right)$$
$$K_{1} := \left(\left[2^{e-m} N_{1} u \right] P_{u}, \left[2^{e-m} g_{u} s_{uv} \right] P_{v} \right),$$
$$K_{2} := \left(\left[2^{e-m} (N_{1} u + g_{u} g_{v} \mu N_{2} \delta_{uv}^{2}) \right] Q_{u}, \left[2^{e-m} g_{u} s_{uv} \right] Q_{v} \right)$$

is a $\zeta^{2^{e^{-m}}}$ -symplectic basis of $(E_u \times E_v)[2^{m+2}]$ adapted to $\Phi := f_m \circ \cdots \circ f_1 : E_u \times E_v \to A_m$ i.e. such that $\ker(\Phi) = \langle [4]K_1, [4]K_2 \rangle$.

Proof. By Lemma 13, we have:

$$\ker(\Phi) = \langle ([2^{e+2-m}N_1u]P_u, [2^{e+2-m}g_us_{uv}]P_v), \\ ([2^{e+2-m}N_1u]Q_u, [2^{e+2-m}g_us_{uv}]Q_v) \rangle,$$

so that $\ker(\Phi) = \langle [4]K_1, [4]K_2 \rangle$ because $2^{e+2-m}\delta_{uv}^2 \equiv 0 \mod 2^{e+2}$. Besides, we clearly have

$$e_{2^{m+2}}(J_1, J_2) = e_{2^{m+2}}(J_1, K_2) = e_{2^{m+2}}(J_2, K_1) = 1$$

and

$$e_{2^{m+2}}(K_1, K_2) = e_{2^{m+2}}(([2^{e-m}N_1u]P_u, [2^{e-m}g_us_{uv}]P_v),$$

$$([2^{e-m}(N_1u + g_ug_v\mu N_2\delta_{uv}^2)]Q_u, [2^{e-m}g_us_{uv}]Q_v))$$

$$= e_{2^{m+2}}([2^{e-m}]P_u, [2^{e-m}]Q_u)^{N_1u(N_1u+g_ug_v\mu N_2\delta_{uv}^2)}$$

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$$\begin{split} &\cdot e_{2^{m+2}}([2^{e-m}]P_v, [2^{e-m}]Q_v)^{g_u^2 s_{uv}^2} \\ &= e_{2^{e+2}}(P_u, Q_u)^{2^{e-m}N_1u(N_1u+g_ug_v\mu N_2\delta_{uv}^2)} \\ &\cdot e_{2^{e+2}}(P_v, Q_v)^{2^{e-m}g_u^2 s_{uv}^2} \\ &= e_{2^{e+2}}(P, Q)^{2^{e-m}(g_uN_1u(N_1u+g_ug_v\mu N_2\delta_{uv}^2)+g_vN_1N_2g_u^2 s_{uv}^2)} \\ &= \zeta^{2^{e-m}(g_uN_1^2u^2+g_vN_1N_2g_u^2(\delta_{uv}^2+s_{uv}^2))} \\ &= \zeta^{2^{e-m}(g_uN_1^2u^2+g_vN_1N_2g_u^2(x_u^2+y_u^2)(x_v^2+y_v^2))} \\ &= \zeta^{2^{e-m}g_uuN_1(uN_1+vN_2)} = \zeta^{2^{2e-m}} = 1 \end{split}$$

Indeed, Eq. (10) implies that $2^m \leq |\delta_{uv}| \leq x_u^2 + y_u^2 + x_v^2 + y_v^2 \leq u + v = O(2^{e/2})$, so that m = O(e/2) and $2^{2e-m} \equiv 0 \mod 2^{e+2}$. Finally,

$$e_{2^{m+2}}(J_1, K_1) = e_{2^{m+2}}(([-2^{e-m}\eta_1\mu\gamma_u]Q_u, 0), ([2^{e-m}N_1u]P_u, [2^{e-m}g_us_{uv}]P_v))$$

$$= e_{2^{e+2}}(Q_u, P_u)^{-2^{e-m}\eta_1\mu\gamma_uN_1u} = e_{2^{e+2}}(P_u, Q_u)^{2^{e-m}\gamma_u}$$

$$= e_{2^{e+2}}(P, Q)^{2^{e-m}\gamma_ug_u} = \zeta^{2^{e-m}},$$

and

$$e_{2^{m+2}}(J_2, K_2) = e_{2^{m+2}}((0, [2^{e-m}\gamma_u\gamma_v\eta_1\eta_2\sigma_{uv}]P_v), \\ ([2^{e-m}(N_1u + g_ug_v\mu N_2\delta_{uv}^2)]Q_u, [2^{e-m}g_us_{uv}]Q_v)) \\ = e_{2^{e+2}}(P_v, Q_v)^{2^{e-m}\gamma_u\gamma_v\eta_1\eta_2\sigma_{uv}g_us_{uv}} \\ = e_{2^{e+2}}(P, Q)^{2^{e-m}g_vN_1N_2\gamma_u\gamma_v\eta_1\eta_2\sigma_{uv}g_us_{uv}} \\ = \zeta^{2^{e-m}},$$

so (J_1, J_2, K_1, K_2) is indeed a $\zeta^{2^{e^{-m}}}$ -symplectic basis of $(E_u \times E_v)[2^{m+2}]$. \Box Lemma 16. The change of basis matrix from \mathscr{B}_1 to $[2^m]\mathscr{B}_2$ is:

$$M_1 := \begin{pmatrix} 0 & 0 & N_1 u & 0 \\ 0 & \gamma_u \gamma_v \eta_1 \eta_2 \sigma_{uv} & g_u s_{uv} & 0 \\ -\eta_1 \mu & 0 & 0 & g_u N_1 u \\ 0 & 0 & 0 & g_u g_v N_1 N_2 s_{uv} \end{pmatrix}.$$

Proof. It follows immediately form Lemma 15 and the fact that $m \ge 1$ so that $4|\delta_{uv}^2$.

B.3 The 4-dimensional gluing

Let $H := F_m \circ \cdots \circ F_1 : E_u^2 \times E_v^2 \to A_m^2$ as in Lemma 13. Since $H = \text{Diag}(\Phi, \Phi) \circ S$ with $S : (R_1, S_1, R_2, S_2) \mapsto (R_1, R_2, S_1, S_2)$, if we assume we have computed Φ in Appendix B.2, we can evaluate H on any point. Images of H are expressed

in product level 2 theta coordinates on A_m^2 , where the theta structure on A_m is associated to the ζ^{2^e} -symplectic basis of $A_m[4]$ induced by Φ , and given by:

$$\mathscr{C}_0 := ([2^m] \Phi(J_1), [2^m] \Phi(J_2), \Phi(K_1), \Phi(K_2)).$$

Hence, the product theta structure on A_m^2 is induced by $\mathscr{C}_0 \times \mathscr{C}_0$, as defined in Eq. (15).

In general, this basis is not adapted to $F_{m+1}: A_m^2 \to B$. Such a basis is given by:

$$\mathscr{C}_1 := ([2^e]H(S_1), \cdots, [2^e]H(S_4), [2^{e-m}]H(T_1), \cdots, [2^{e-m}]H(T_4)),$$

where $\mathscr{C} := (S_1, \cdots, S_4, T_1, \cdots, T_4)$ is a ζ -symplectic basis of $(E_u^2 \times E_v^2)[2^{e+2}]$ adapted to F. Applying Lemma 12 to $(x_1, x_2, y_1, y_2) := ((\varphi_{\mathfrak{b}_k}(P), 0), (0, \varphi_{\mathfrak{b}_k}(P)), ([\eta_1]\varphi_{\mathfrak{b}_k}(Q), 0), (0, [\eta_1]\varphi_{\mathfrak{b}_k}(Q)))$, we obtain:

$$\begin{split} S_1 &:= \left([-\mu\eta_1 x_u] Q_u, [-\mu\eta_1 y_u] Q_u, 0, 0 \right), \quad S_2 := \left([\mu\eta_1 y_u] Q_u, [-\mu\eta_1 x_u] Q_u, 0, 0 \right), \\ S_3 &:= \left(0, 0, [\nu\eta_2 x_v] P_v, [\nu\eta_2 y_v] P_v \right), \quad S_4 := \left(0, 0, [-\nu\eta_2 y_v] P_v, [\nu\eta_2 x_v] P_v \right), \\ T_1 &:= \left([N_1 x_u] P_u, [N_1 y_u] P_u, [x_v] P_v, [y_v] P_v \right), \\ T_2 &:= \left([-N_1 y_u] P_u, [N_1 x_u] P_u, [-y_v] P_v, [x_v] P_v \right), \\ T_3 &:= \left([(1 - 2^e \mu \eta_1) x_u] Q_u, [(1 - 2^e \mu \eta_1) y_u] Q_u, [\eta_1 x_v] Q_v, [\eta_1 y_v] Q_v \right), \\ T_4 &:= \left([-(1 - 2^e \mu \eta_1) y_u] Q_u, [(1 - 2^e \mu \eta_1) x_u] Q_u, [-\eta_1 y_v] Q_v, [\eta_1 x_v] Q_v \right), \end{split}$$

where μ, ν, η_1, η_2 are inverses of u, v, N_1, N_2 modulo 2^{e+2} respectively.

To compute F_{m+1} , we need to compute the symplectic change of basis from $\mathscr{C}_0 \times \mathscr{C}_0$ to \mathscr{C}_1 and the associated change of theta coordinates. We can then apply the algorithms from [18, § 4.3] to compute F_{m+1} with level 2 theta coordinates.

Lemma 17. The change of basis matrix from $\mathscr{C}_0 \times \mathscr{C}_0$ to \mathscr{C}_1 is

$$M_3 := \begin{pmatrix} g_u x_u - g_u y_u & 0 & 0 & 0 & \mu_2 & \mu_3 \\ 0 & 0 & \lambda x_v - \lambda y_v & \mu_1 y_u & \mu_1 x_u & 0 & 0 \\ g_u y_u & g_u x_u & 0 & 0 & 0 & 0 & -\mu_3 & \mu_2 \\ 0 & 0 & \lambda y_v & \lambda x_v & -\mu_1 x_u & \mu_1 y_u & 0 & 0 \\ 0 & 0 & 0 & 0 & \mu x_u & -\mu y_u & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & \eta_1 x_v \gamma_u \sigma_{uv} -\eta_1 y_v \gamma_u \sigma_{uv} \\ 0 & 0 & 0 & 0 & \mu y_u & \mu x_u & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & \eta_1 y_v \gamma_u \sigma_{uv} & \eta_1 x_v \gamma_u \sigma_{uv} \end{pmatrix},$$

where $\lambda := \nu g_u g_v N_1 s_{uv}$, $\mu_1 := \mu g_u^2 g_v s_{uv} N_1 N_2 \delta_{uv}/2^m$,

$$\mu_2 := \frac{\delta_{uv}}{2^m} g_u \sigma_{uv} (N_1 u y_u + g_v x_v N_2 \delta_{uv}) \quad and$$
$$\mu_3 := \frac{\delta_{uv}}{2^m} g_u \sigma_{uv} (N_1 u x_u - g_v y_v N_2 \delta_{uv}).$$

Proof. We have:

$$\begin{aligned} [2^e]H(S_1) &= \left(\varPhi([-2^e \mu \eta_1 x_u] Q_u, 0), \varPhi([-2^e \mu \eta_1 y_u] Q_u, 0) \right) \\ &= \left([g_u x_u] \varPhi([-2^e \eta_1 \mu \gamma_u] Q_u, 0), [g_u y_u] \varPhi([-2^e \eta_1 \mu \gamma_u] Q_u, 0) \right) \\ &= \left([g_u x_u] [2^m] \varPhi(J_1), [g_u y_u] [2^m] \varPhi(J_1) \right) \end{aligned}$$

$$\begin{split} [2^e]H(S_2) &= (\varPhi([2^e\mu\eta_1 y_u]Q_u, 0), \varPhi([-2^e\mu\eta_1 x_u]Q_u, 0)) \\ &= ([-g_u y_u][2^m]\varPhi(J_1), [g_u x_u][2^m]\varPhi(J_1)) \end{split}$$

$$\begin{split} [2^e]H(S_3) &= (\varPhi(0, [2^e \nu \eta_2 x_v] P_v), \varPhi(0, [2^e \nu \eta_2 y_v] P_v)) \\ &= ([\nu x_v g_u g_v N_1 s_{uv}] \varPhi(0, [2^e \gamma_u \gamma_v \eta_1 \eta_2 \sigma_{uv}] P_v), \\ & [\nu y_v g_u g_v N_1 s_{uv}] \varPhi(0, [2^e \gamma_u \gamma_v \eta_1 \eta_2 \sigma_{uv}] P_v)) \\ &= ([\lambda x_v] [2^m] \varPhi(J_2), [\lambda y_v] [2^m] \varPhi(J_2)) \end{split}$$

$$[2^{e}]H(S_{4}) = (\Phi(0, [-2^{e}\nu\eta_{2}y_{v}]P_{v}), \Phi(0, [2^{e}\nu\eta_{2}x_{v}]P_{v}) = ([-\lambda y_{v}][2^{m}]\Phi(J_{2}), [\lambda x_{v}][2^{m}]\Phi(J_{2})).$$

We also have:

$$[2^{e-m}]H(T_1) = (\Phi([2^{e-m}N_1x_u]P_u, [2^{e-m}x_v]P_v), \Phi([2^{e-m}N_1y_u]P_u, [2^{e-m}y_v]P_v)),$$

with:

$$\begin{split} \varPhi([2^{e-m}N_1x_u]P_u, [2^{e-m}x_v]P_v) &= [\mu x_u]\varPhi([2^{e-m}N_1u]P_u, [2^{e-m}g_us_{uv}]P_v) \\ &+ \varPhi(0, [2^{e-m}(x_v - \mu x_ug_us_{uv})]P_v) \\ &= [\mu x_u]\varPhi(K_1) + \varPhi(0, [2^{e-m}\mu g_uy_u(x_vy_u - x_uy_v)]P_v) \\ &= [\mu x_u]\varPhi(K_1) \\ &+ [\mu g_u^2g_vy_u\delta_{uv}s_{uv}N_1N_2]\varPhi(0, [2^{e-m}\gamma_u\gamma_v\eta_1\eta_2\sigma_{uv}]P_v) \\ &= [\mu_1y_u][2^m]\varPhi(J_2) + [\mu x_u]\varPhi(K_1) \end{split}$$

and:

$$\begin{split} \varPhi([2^{e-m}N_1y_u]P_u, [2^{e-m}y_v]P_v) &= [\mu y_u]\varPhi(K_1) \\ &+ \varPhi(0, [2^{e-m}(y_v - \mu y_u g_u s_{uv})]P_v) \\ &= [\mu y_u]\varPhi(K_1) + \varPhi(0, [-2^{e-m}\mu g_u x_u(x_v y_u - x_u y_v)]P_v) \\ &= [\mu y_u]\varPhi(K_1) \\ &- [\mu g_u^2 g_v x_u \delta_{uv} s_{uv} N_1 N_2]\varPhi(0, [2^{e-m}\gamma_u \gamma_v \eta_1 \eta_2 \sigma_{uv}]P_v) \\ &= -[\mu_1 x_u][2^m]\varPhi(J_2) + [\mu y_u]\varPhi(K_1), \end{split}$$

so that:

$$[2^{e-m}]H(T_1) = ([\mu_1 y_u][2^m] \varPhi(J_2) + [\mu x_u] \varPhi(K_1), -[\mu_1 x_u][2^m] \varPhi(J_2) + [\mu y_u] \varPhi(K_1))$$

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And we also have

$$[2^{e-m}]H(T_2) = (\Phi([-2^{e-m}N_1y_u]P_u, [-2^{e-m}y_v]P_v), \Phi([2^{e-m}N_1x_u]P_u, [2^{e-m}x_v]P_v))$$

= $([\mu_1x_u][2^m]\Phi(J_2) - [\mu y_u]\Phi(K_1), [\mu_1y_u][2^m]\Phi(J_2) + [\mu x_u]\Phi(K_1))$

Similarly, we compute

$$[2^{e-m}]H(T_3) = (T_{3,1}, T_{3,2}),$$

with:

$$\begin{split} T_{3,1} &:= \varPhi([2^{e-m}(1-2^e\mu\eta_1)x_u]Q_u, [2^{e-m}\eta_1x_v]Q_v) \\ &= \varPhi([2^{e-m}x_u]Q_u, [2^{e-m}\eta_1x_v]Q_v) \\ &= [\eta_1x_v\gamma_u\sigma_{uv}]\varPhi([2^{e-m}(N_1u+g_ug_v\mu N_2\delta_{uv}^2)]Q_u, [2^{e-m}g_us_{uv}]Q_v) \\ &+ \varPhi([2^{e-m}(x_u-\eta_1x_v\gamma_u\sigma_{uv}(N_1u+g_ug_v\mu N_2\delta_{uv}^2))]Q_u, 0) \\ &= [\eta_1x_v\gamma_u\sigma_{uv}]\varPhi(K_2) \\ &+ \varPhi([-2^{e-m}\delta_{uv}\eta_1\sigma_{uv}\mu(N_1uy_u+g_vx_vN_2\delta_{uv})]Q_u, 0) \\ &= [\eta_1x_v\gamma_u\sigma_{uv}]\varPhi(K_2) \\ &+ [\delta_{uv}g_u\sigma_{uv}(N_1uy_u+g_vx_vN_2\delta_{uv})]\varPhi([-2^{e-m}\eta_1\mu\gamma_u]Q_u, 0) \\ &= [\mu_2][2^m]\varPhi(J_1) + [\eta_1x_v\gamma_u\sigma_{uv}]\varPhi(K_2) \end{split}$$

and:

$$\begin{split} T_{3,2} &:= \varPhi([2^{e-m}(1-2^e\mu\eta_1)y_u]Q_u, [2^{e-m}\eta_1y_v]Q_v) \\ &= \varPhi([2^{e-m}y_u]Q_u, [2^{e-m}\eta_1y_v]Q_v) \\ &= [\eta_1y_v\gamma_u\sigma_{uv}]\varPhi([2^{e-m}(N_1u+g_ug_v\mu N_2\delta_{uv}^2)]Q_u, [2^{e-m}g_us_{uv}]Q_v) \\ &+ \varPhi([2^{e-m}(y_u-\eta_1y_v\gamma_u\sigma_{uv}(N_1u+g_ug_v\mu N_2\delta_{uv}^2)]Q_u, 0) \\ &= [\eta_1y_v\gamma_u\sigma_{uv}]\varPhi(K_2) \\ &+ \varPhi([2^{e-m}\delta_{uv}\eta_1\sigma_{uv}\mu(N_1ux_u-y_vg_ug_vN_2\delta_{uv})]Q_u, 0) \\ &= [\eta_1y_v\gamma_u\sigma_{uv}]\varPhi(K_2) \\ &- [\delta_{uv}g_u\sigma_{uv}(N_1ux_u-g_ug_vy_vN_2\delta_{uv})]\varPhi([-2^{e-m}\eta_1\mu\gamma_u]Q_u, 0) \\ &= -[\mu_3]\varPhi(J_1) + [\eta_1y_v\gamma_u\sigma_{uv}]\varPhi(K_2), \end{split}$$

so that:

$$[2^{e-m}]H(T_3) = ([\mu_2][2^m]\Phi(J_1) + [\eta_1 x_v \gamma_u \sigma_{uv}]\Phi(K_2), - [\mu_3][2^m]\Phi(J_1) + [\eta_1 y_v \gamma_u \sigma_{uv}]\Phi(K_2)).$$

Finally, we have

$$\begin{split} [2^{e-m}]H(T_4) &= (\varPhi([-(1-2^e\mu\eta_1)y_u]Q_u, [-\eta_1y_v]Q_v), \\ & \varPhi([(1-2^e\mu\eta_1)x_u]Q_u, [\eta_1x_v]Q_v)) = (-T_{3,2}, T_{3,1}) \\ &= ([\mu_3][2^m]\varPhi(J_1) - [\eta_1y_v\gamma_u\sigma_{uv}]\varPhi(K_2), \\ & [\mu_2][2^m]\varPhi(J_1) + [\eta_1x_v\gamma_u\sigma_{uv}]\varPhi(K_2)). \end{split}$$

This completes the proof.

B.4 The splitting and identification of the result curve $E_{\mathfrak{a}}$

Once we have computed $F : E_u^2 \times E_v^2 \to E_a^2 \times E'^2$ as a 2-isogeny chain, the codomain we obtain does not have a product theta structure from which we can obtain the theta null point of E_a , using Eq. (16). The theta structure we obtain is induced by the ζ^{2^e} -symplectic basis of $(E_a^2 \times E'^2)[4]$ given by:

 $\mathscr{D}_1 := ([2^e]F(S_1), \cdots, [2^e]F(S_4), F(T_1), \cdots, F(T_4)),$

while a product theta structure of $E_{\mathfrak{a}}^2 \times E'^2$ is induced by the ζ^{2^e} -symplectic basis $\mathscr{D}_0 := (U_1, \cdots, U_4, V_1, \cdots, V_4)$, where:

$$U_{1} := ([2^{e}]\varphi_{\mathfrak{b}_{k}}(P), 0, 0, 0), \quad U_{2} := (0, [2^{e}]\varphi_{\mathfrak{b}_{k}}(P), 0, 0),$$
$$U_{3} := (0, 0, [2^{e}]\Psi \circ \widetilde{\Phi}(\varphi_{\mathfrak{b}_{k}}(P), 0)), \quad U_{4} := (0, 0, [2^{e}]\Psi \circ \widetilde{\Phi}(0, \varphi_{\mathfrak{b}_{k}}(P))),$$
$$V_{1} := ([2^{e}\eta_{1}]\varphi_{\mathfrak{b}_{k}}(Q), 0, 0, 0), \quad V_{2} := (0, [2^{e}\eta_{1}]\varphi_{\mathfrak{b}_{k}}(Q), 0, 0),$$

 $V_3 := (0, 0, [2^e \mu \nu \eta_1^2 \eta_2] \Psi \circ \tilde{\varPhi}(\varphi_{\mathfrak{b}_k}(Q), 0)), \quad V_4 := (0, 0, [2^e \mu \nu \eta_1^2 \eta_2] \Psi \circ \tilde{\varPhi}(0, \varphi_{\mathfrak{b}_k}(Q))),$

given by Lemma 12, where η_1, η_2, μ, ν are modular inverse of N_1, N_2, u, v modulo 4 respectively. By Lemma 12, we also obtain the

Lemma 18. The change of basis matrix from \mathscr{D}_1 to \mathscr{D}_0 is given by

$$M_4 := \begin{pmatrix} 0 & 0 & 0 & 0 & -1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & -1 & 0 & 0 \\ 0 & 0 & vN_2 & 0 & 0 & 0 & 0 & 0 \\ 0 & 0 & vN_2 & 0 & 0 & 0 & 0 & 0 \\ 1 & 0 & -vN_2 & 0 & 0 & 0 & 0 & 0 \\ 0 & 1 & 0 & -vN_2 & 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & \nu\eta_2 & 0 \\ 0 & 0 & 0 & 0 & 1 & 0 & \nu\eta_2 \end{pmatrix}$$

Hence, using the matrix M_4 above, the non-product theta null point of $E_{\mathfrak{a}}^2 \times E'^2$ and the change of coordinates formulas from [18, Theorem 12], we can compute the product theta null point of $E_{\mathfrak{a}}^2 \times E'^2$ given by:

$$(\theta_{i_1,i_2,i_3,i_4}^{E_{\mathfrak{a}}^2 \times E'^2}(0))_{i_1,i_2,i_3,i_4 \in \mathbb{Z}/2\mathbb{Z}} = (\theta_{i_1}^{E_{\mathfrak{a}}}(0) \cdot \theta_{i_2}^{E_{\mathfrak{a}}}(0) \cdot \theta_{i_3}^{E'}(0) \cdot \theta_{i_4}^{E'}(0))_{i_1,i_2,i_3,i_4 \in \mathbb{Z}/2\mathbb{Z}}.$$
 (17)

We can extract the projective theta null point $(a:b) := (\theta_0^{E_{\mathfrak{a}}}(0): \theta_1^{E_{\mathfrak{a}}}(0))$ by finding $i_3, i_4 \in \mathbb{Z}/2\mathbb{Z}$ such that the $\theta_{0,0,i_3,i_4}^{E_{\mathfrak{a}}^2 \times E'^2}(0) \neq 0$ and set $a := \theta_{0,0,i_3,i_4}^{E_{\mathfrak{a}}^2 \times E'^2}(0)$ and $b := \theta_{1,0,i_3,i_4}^{E_{\mathfrak{a}}^2 \times E'^2}(0)$. This theta null point (a:b) give 6 different possibilities for the Montgomery coefficient A of

$$E_{\mathfrak{a}}: By^2 = x^3 + Ax^2 + x.$$

Those possibilities are presented in Proposition 11. All of these possible coefficients are not defined over \mathbb{F}_p as $E_{\mathfrak{a}}$ should. This removes 2 to 4 possibilities out of 6.

However, we still cannot distinguish $E_{\mathfrak{a}}$ from its quadratic twist *i.e.* identify B from the theta null point (a : b). To circumvent this difficulty, we impose B = 1 (or any quadratic residue over \mathbb{F}_p) and require that 4-torsion points of the form $(\pm 1 : * : 1)$ are defined in $E_{\mathfrak{a}}(\mathbb{F}_p)$ *i.e.* that A+2 and A-2 are quadratic residues over \mathbb{F}_p . This lifts all ambiguity since a Montgomery curve is determined by a point of 4-torsion.

In practice, to identify the right Montgomery coefficient A of $E_{\mathfrak{a}}$, we pick a 4-torsion point $T_u \in E_u(\mathbb{F}_p)$, which does exist since E_u is defined over \mathbb{F}_p , as the result of an ideal action on E (see the proof of Lemma 5.1). Since

$$F(T_u, 0, 0, 0) = ([x_u]\varphi_{\mathfrak{b}_k} \circ \widehat{\varphi}_u(T_u), *),$$

and both $\varphi_{\mathfrak{b}_k}$ and φ_u are associated to ideal actions, the point $T_{\mathfrak{a}} := [x_u]\varphi_{\mathfrak{b}_k} \circ \widehat{\varphi}_u(T_u)$ is \mathbb{F}_p -rational. Because $N_1 = N(\mathfrak{a})$ and $g_u = \deg(\varphi_u)$ are odd and x_u can be assumed to be odd (swapping x_u and y_u if necessary), $T_{\mathfrak{a}}$ is also a point of 4-torsion. We can then compute several change of theta coordinates until $T_{\mathfrak{a}}$ is mapped to one of the \mathbb{F}_p -rational theta points (1:0) or (0:1) that are mapped to the desired \mathbb{F}_p -rational Montgomery points (-1:1) and (1:1) respectively. Once this is done, we identify a Montgomery coefficient A form the new theta null point, following Proposition 11.

To lift $(\pm 1 : 1)$ in $E_{\mathfrak{a}}(\mathbb{F}_p)$ given by $y^2 = x^3 + Ax^2 + x$, we need $A \pm 2$ to be a quadratic residue over \mathbb{F}_p . If this is not the case, then $E_{\mathfrak{a}}$ is given by $-y^2 = x^3 + Ax^2 + x$ (in which $(\pm 1 : 1)$ is \mathbb{F}_p -rational). But there is an \mathbb{F}_p -rational isomorphism $(x, y) \mapsto (-x, y)$ from $-y^2 = x^3 + Ax^2 + x$ to $y^2 = x^3 - Ax^2 + x$ so we can change A into -A in this case.

In practice, on the Montgomery curve E_u , we take T_u the point with xcoordinate $x(T_u) = 1$. By the same reasoning as above, replacing A_u by $-A_u$ if necessary, we can assume that T_u is rational. The level 2 theta null point we obtain in the end for E_a implicitly contains the x-coordinate of the image of T_u , so we do not even need to push the 4-torsion point T_u explicitly.

C Relaxing the **B**-good condition

This section outlines an approach to relax the \mathfrak{B} -good condition of Section 4.1 to improve the probability of finding (u, v) such that isogenies of degree u and v can be efficiently computed in dimension 2.

Instead of looking for a pair (u, v) of \mathfrak{B} -good integers, we shall search for (u, v) such that u takes the form $u = g_u(ax_u^2 + by_u^2)$ and v the form $v = g_v(ax_v^2 + by_v^2)$, where $x_u, y_u, x_v, y_v \in \mathbb{N}$ and $g_u, g_v, a, b \in \mathbb{N}^*$ are products of primes lying in \mathfrak{B} . Note that the \mathfrak{B} -good case is the case a = b = 1.

Before discussing the advantages and drawbacks of introducing additional freedom through a and b, let us describe how the Kani diagram in dimension 4 for computing $E_{\mathfrak{a}}$ (cf. Section 3.1) is reshaped by this change. In this construction, we assume that \mathfrak{B} is a set of Elkies primes. As we shall see, this assumption is crucial to ensure the well-definedness of our Kani diagram.

We have $uN(\mathfrak{b}_k) + vN(\mathfrak{c}_k) = 2^e$ where $\mathfrak{b} = \mathfrak{b}_e \mathfrak{b}_k$ and $\mathfrak{c} = \mathfrak{c}_e \mathfrak{c}_k$ are ideals equivalent to \mathfrak{a} with $[\mathfrak{b}_k] \cdot E_1 = E_{\mathfrak{a}}$ and $[\mathfrak{c}_k] \cdot E_2 = E_{\mathfrak{a}}$. Additionally, $u = g_u(ax_u^2 + by_u^2)$ and $v = g_v(ax_v^2 + by_v^2)$ as defined above.

Using Elkies' algorithm (cf. Section 3.2), we construct the following (ax_u^2, by_u^2) -isogeny diamond and (ax_v^2, by_v^2) -isogeny diamond

$$\begin{array}{ccc} E_{1,b} \xrightarrow{x_u \varphi'_{1,a}} E_{1,ab} & E_{2,b} \xrightarrow{x_v \varphi'_{2,a}} E_{2,ab} \\ y_u \varphi_{1,b} & & & & & & \\ y_u \varphi_{1,b} & & & & & & \\ & & & & & & \\ E_1 \xrightarrow{x_u \varphi_{1,a}} E_{1,a} & & & & & E_2 \xrightarrow{x_v \varphi_{2,a}} E_{2,a} \end{array}$$

where $\varphi_{1,a}, \varphi'_{1,a}, \varphi_{2,a}, \varphi'_{2,a}$ are isogenies of degree a and $\varphi_{1,b}, \varphi'_{1,b}, \varphi_{2,b}, \varphi'_{2,b}$ are isogenies of degree b. From this, we obtain an $(ax_u^2 + by_u^2)$ -isogeny $\Phi_1 : E_1 \times E_{1,ab} \to E_{1,a} \times E_{1,b}$ and an $(ax_v^2 + by_v^2)$ -isogeny $\Phi_2 : E_2 \times E_{2,ab} \to E_{2,a} \times E_{2,b}$ explicitly given by the matrices

$$\begin{pmatrix} x_u\varphi_{1,a} & y_u\hat{\varphi}'_{1,b} \\ -y_u\varphi_{1,b} & x_u\hat{\varphi}'_{1,a} \end{pmatrix} \text{ and } \begin{pmatrix} x_v\varphi_{2,a} & y_v\hat{\varphi}'_{2,b} \\ -y_v\varphi_{2,b} & x_v\hat{\varphi}'_{2,a} \end{pmatrix}$$

Then, we compute a g_u -isogeny $\Phi_{g_u} : E_{1,a} \times E_{1,b} \to A_u$ and a g_v -isogeny $\Phi_{g_v} : E_{2,a} \times E_{2,b} \to A_v$, where A_u and A_v are each products of elliptic curves. This can be achieved using Elkies' algorithm to compute four isogenies $\psi_{a,u}, \psi_{b,u}, \psi_{a,v}$ and $\psi_{b,v}$ such that $\psi_{c,u} : E_{1,c} \to E_{c,u}$ is an g_u -isogeny and $\psi_{c,v} : E_{1,c} \to E_{c,v}$ is an g_v -isogeny, where $c \in \{a, b\}$. Hence, we obtain Φ_{g_u} and Φ_{g_v} from the diagonal matrices

$$\begin{pmatrix} \psi_{a,u} & 0\\ 0 & \psi_{b,u} \end{pmatrix} \text{ and } \begin{pmatrix} \psi_{a,v} & 0\\ 0 & \psi_{b,v} \end{pmatrix}.$$

Let us denote by Φ_u the $g_u(ax_u^2 + by_u^2)$ -isogeny $\Phi_{g_u} \circ \Phi_1 : E_1 \times E_{1,ab} \to A_u$ and by Φ_v the $g_v(ax_v^2 + by_v^2)$ -isogeny $\Phi_{g_v} \circ \Phi_2 : E_2 \times E_{2,ab} \to A_v$.

To state the Kani diagram in dimension 4, it remains only to consider the actions of \mathbf{b}_k and \mathbf{c}_k . Note that the key difference here is that \mathbf{b}_k (resp. \mathbf{c}_k) acts on $E_1 \times E_{1,ab}$ (resp. $E_2 \times E_{2,ab}$) rather than on E_1^2 (resp. E_2^2). We denote by $\Phi_{\mathbf{b}_k} : E_1 \times E_{1,ab} \to E_{\mathfrak{a}} \times E_3$ and $\Phi_{\mathbf{c}_k} : E_2 \times E_{2,ab} \to E_{\mathfrak{a}} \times E_4$ the corresponding $N(\mathbf{b}_k)$ -isogeny and $N(\mathbf{c}_k)$ -isogeny. To ensure that this leads to a construction similar to the \mathfrak{B} -good case, we now prove that $E_3 = E_4$.

Thanks to the assumption on the primes in \mathfrak{B} being Elkies, we can express E_3 and E_4 in terms of ideal actions. We have

$$E_3 = [\mathfrak{b}_k] \cdot E_{1,ab} = [\mathfrak{b}_k I_a I_b] \cdot E_1 = [\mathfrak{b}_k I_a I_b \mathfrak{b}_e] \cdot E$$

and

$$E_4 = [\mathfrak{c}_k] \cdot E_{2,ab} = [\mathfrak{c}_k I_a I_b] \cdot E_2 = [\mathfrak{c}_k I_a I_b \mathfrak{c}_e] \cdot E_2$$

By commutativity and by construction of \mathfrak{b}_k and \mathfrak{c}_k , we have

$$[\mathfrak{b}_k I_a I_b \mathfrak{b}_e] = [I_a I_b \mathfrak{b}] \text{ and } [\mathfrak{c}_k I_a I_b \mathfrak{c}_e] = [I_a I_b \mathfrak{c}].$$

Moreover, since both \mathfrak{b} and \mathfrak{c} are equivalent to \mathfrak{a} , we have that $E_3 = [I_a I_b] \cdot E_{\mathfrak{a}} = E_4$. We denote by E' the curve $[I_a I_b] \cdot E_{\mathfrak{a}}$.

Finally, using Lemma 3, there exist a $g_u(ax_u^2 + by_u^2)N(\mathfrak{c}_k)$ -isogeny $\Psi: A_u \to A$ and a $g_v(ax_v^2 + by_v^2)N(\mathfrak{b}_k)$ -isogeny $\Phi: A \to A_v$ such that $\Phi \circ \Psi = \Phi_v \circ \tilde{\Phi}_{\mathfrak{c}_k} \circ \Phi_{\mathfrak{b}_k} \circ \tilde{\Phi}_u$. Therefore, we have the following Kani-square



By Lemma 4 and as $g_u(ax_u^2 + by_u^2)N_1 + g_v(ax_v^2 + by_v^2)N_2 = 2^e$, this isogenydiamond leads to an 2^e -isogeny $F: A_u \times A_v \to E_{\mathfrak{a}} \times E' \times A$ with kernel

$$\ker(F) = \{ ([g_u(ax_u^2 + by_u^2)N_1](x), \Phi_v \circ \widetilde{\Phi}_{\mathfrak{c}_k} \circ \Phi_{\mathfrak{b}_k} \circ \widetilde{\Phi}_u(x) | x \in A_u[2^e] \}.$$

This kernel can also be expressed from points of $E_1 \times E_{1,ab}$ as

$$\ker(F) = \{ ([N_1]\Phi_u(P,Q), \Phi_v \circ \widetilde{\Phi}_{\mathfrak{c}_k} \circ \Phi_{\mathfrak{b}_k}(P,Q)) | (P,Q) \in E_1 \times E_{1,ab}[2^e] \}.$$

From there, one can compute this 2^e -isogeny F using the same method as in Section 3.1.

Assessment of the relaxation. We now investigate the benefits and drawbacks of considering pairs (u, v) of such a form. First, the main additional cost is the computation of the *a*-isogenies $\varphi_{1,a}, \varphi'_{1,a}, \varphi_{2,a}, \varphi'_{2,a}$ and *b*-isogenies $\varphi_{1,b}, \varphi'_{1,b}, \varphi_{2,b}, \varphi'_{2,b}$. Since half of them are simply pushforwards of the other half, they can essentially be computed with two calls to the Elkies' algorithm for the *a*-isogenies and two calls for the *b*-isogenies. On the other hand, one may expect that the probability of an integer being representable as $ax^2 + by^2$, with $a, b \in \{1, 2, 3, 5, 7\}$, is at least three times greater than the probability of being representable by a sum of two squares. The table of experimental results in Figure 1 supports this assumption.

Let us briefly explain why the set $\{2, 3, 5, 7\}$ is an interesting choice for \mathfrak{B} regarding a, b.

We first focus on forms $x^2 + by^2$ where b is prime. Our discussion relies mainly on Proposition C.1 and on the fact that an integer whose prime factors are all represented by such a quadratic form can itself be represented by this form.

Proposition C.1. [17, Corollary 2.6] Let b be an integer and p be an odd prime such that gcd(b,p) = 1. Then p is represented by a primitive form of discriminant -4b if and only if its Legendre symbol (-b/p) is equal to 1.

Integer length	32 bits	64 bits	128 bits
Success rate when $a, b \in \{1\}$	0.11	0.08	0.05
Success rate when $a, b \in \{1, 2, 3, 7\}$	0.31	0.24	0.18
Success rate when $a, b \in \{1, 2, 3, 5, 7\}$	0.33	0.28	0.20

Fig. 1. Success rates for representing an integer as $ax^2 + by^2$, with $x, y \in \mathbb{N}$, depending on the possible values for a and b. Each rate is computed from 10,000 integers drawn uniformly at random. Forms where $a \neq 1$ and b = 7 were not considered.

Note that the quadratic form $x^2 + by^2$ has discriminant -4b. Hence, the best choices for b to maximize the probability that primes are represented by $x^2 + by^2$ are those for which the class group of discriminant -4b has cardinality 1. Indeed, by Proposition C.1, if (b/p) = 1, then p is represented by a primitive form of discriminant 4b. When there is a unique class in this group, we have that $x^2 + by^2$ is equivalent to this primitive form. Therefore, $x^2 + by^2$ also represents p.

Since an integer has a probability of one-half of being a quadratic residue modulo a prime p, we expect that $x^2 + by^2$ represents p with probability one-half. In addition, since the set of even discriminants of class number 1 is -4, -8, -12, -16, -28, there are only 4 such forms interesting to us: $x^2 + y^2$, $x^2 + 2y^2$, $x^2 + 3y^2$ and $x^2 + 7y^2$. We cover all of them with the set of primes $\{2, 3, 7\}$.

The second-best choices for b are those for which the corresponding class group has cardinality 2, such as b = 5 and b = 13. The probability that a prime is representable by $x^2 + by^2$ is then halved for such b. However, from [17], we have the following equivalences

 $\begin{array}{ll} p = x^2 + y^2 & \mbox{iff } p = 1 \mod 4, \\ p = x^2 + 5y^2 & \mbox{iff } p = 1,9 \mod 20, \\ p = x^2 + 13y^2 & \mbox{iff } p = 1,9,17,25,29,49 \mod 52. \end{array}$

Hence, if p is represented by $x^2 + 5y^2$ or $x^2 + 13y^2$, it is also represented by $x^2 + y^2$. Nevertheless, adding 5 to the list of prime factors gives access to the forms $2x^2 + 5x^2$, $x^2 + 10x^2$ and so on. This explains why, in Fig. 1, adding 5 improves the probability of success.

Other experiments have shown that adding a quadratic form from a class group with cardinality 3 or more, such as $x^2 + 11y^2$, has only a negligible impact. This is why, we did not consider forms $ax^2 + by^2$ where $a \neq 1$ and b = 7, nor forms where a or b are composite numbers. All these cases lead to class groups of cardinality greater than 3. Additionally, the next prime after 13 that yields a class group of cardinality 2 is 37. Therefore, to achieve the most significant improvement with the smallest set of primes, it seems reasonable to consider only primes up to 7.

D Algorithms for working over \mathbb{F}_p

In this appendix, we give the necessary algorithms for working over \mathbb{F}_p .

D.1 Fast sampling of a basis given by the eigenspaces

Let $p \equiv 7 \pmod{8}$, and let E/\mathbb{F}_p be a supersingular elliptic curve on the surface, *i.e.* oriented by $\mathbb{Z}[\frac{1+\sqrt{-p}}{2}]$. Recall that for $2^f \parallel (p+1)$, we have

$$E(\mathbb{F}_p)[2^f] = \mathbb{Z}/2\mathbb{Z} \times \mathbb{Z}/2^{f-1}\mathbb{Z},$$

and that on $E[2^{f-1}]$, the frobenius has eigenvalues ± 1 . Our goal is to efficiently sample a basis $\langle P, Q \rangle = E[2^{f-1}]$, such that $x(P), x(Q) \in \mathbb{F}_p$. To do this, we first study the three non-trivial 2-torsion points on E. Denote them as follows:

- T_0 is the point corresponding to the descending 2-isogeny, *i.e.* $E/\langle T_0 \rangle$ is primitively oriented by $\mathbb{Z}[\sqrt{-p}]$
- $-T_{-1}$ is the point corresponding to the *horizontal* 2-isogeny, such that there does not exist a point $T' \in E(\mathbb{F}_p)$ satisfying $[2]T' = T_{-1}$.
- T_1 is the point corresponding to the *horizontal* 2-isogeny, such that there does exist a point $T' \in E(\mathbb{F}_p)$ satisfying $[2]T' = T_1$.

Although these points could easily be distinguished by looking at their division points, we show that they can more efficiently be distinguished by their Tate-pairings.

Lemma D.1. With the notation above, the (reduced) Tate-pairings are given as follows:

 $\begin{array}{l} - \ e_{T,2}(T_{-1},T_{-1}) = e_{T,2}(T_{-1},T_0) = e_{T,2}(T_{-1},T_1) = 1, \\ - \ e_{T,2}(T_0,T_{-1}) = e_{T,2}(T_0,T_0) = -1 \ and \ e_{T,2}(T_0,T_1) = 1, \\ - \ e_{T,2}(T_1,T_{-1}) = e_{T,2}(T_1,T_0) = -1, \ and \ e_{T,2}(T_1,T_1) = 1. \end{array}$

Proof. Recall (see *e.g.* [56, 59, Lemma 5]) that for T_i and $\phi_i : E \to E/\langle T_i \rangle$, we have that $e_{T,2}(T_i, R)$ is 1 if there exists a rational point R' such that $\hat{\phi}_i(R') = R$, and -1 otherwise. The lemma can then be proved as follows:

Note first that we certainly have $e_{T,2}(T_i, T_1) = 1$, since there exists a point T' such that $[2]T' = \hat{\phi}_i \circ \phi_i(T') = T_1$.

For T_0 , the corresponding isogeny ϕ_0 is descending, and hence the 2^f -torsion on $E/\langle T_0 \rangle$ is cyclic. Thus, T_0 is not in the rational image of $\hat{\phi}_0$, and since we know that $e_{T,2}(T_0, T_1) = 1$, we get that $e_{T,2}(T_0, T_0) = e_{T,2}(T_0, T_{-1}) = -1$.

For T_{-1} , we have that ker $(\phi_{-1}) = \langle \phi_1(T_1) \rangle$. Since $f \geq 3$, there are rational points $R' \in E/\langle T_{-1} \rangle$ (of order 4) satisfying $\phi_{-1}(R') = T_1$, and since $E/\langle T_{-1} \rangle$ [2] is \mathbb{F}_p -rational, the other 2-torsion points are mapped to T_{-1} and T_0 . Hence $e_{T,2}(T_{-1},T_i) = 1$.

Finally, for T_1 , we have that $\ker(\widehat{\phi}_1) = \langle \phi_1(T_0) \rangle = \langle \phi_1(T_{-1}) \rangle$. Since T_0 and T_{-1} are both in $E(\mathbb{F}_p) \setminus [2] E(\mathbb{F}_p)$, its clear that $e_{T,2}(T_1, T_0) = e_{T,2}(T_1, T_{-1}) = -1$. (This also follows by the non degeneracy of the Tate pairing and the fact that $e_{T,2}(T_1, T_1) = 1$.)

Thus, given a curve and the three non-trivial 2-torsion points, we can by Lemma D.1 easily detect which point corresponds to which index. Together with the following lemma, this gives a particularily fast algorithm for sampling a basis of the form that we want.

Lemma D.2. Let $p \equiv 7 \pmod{8}$, let

$$E/\mathbb{F}_p: y^2 = g(x)$$

be an elliptic curve with $\operatorname{End}_{\mathbb{F}_p}(E) = \mathbb{Z}[\frac{1+\pi}{2}]$, and let $f = v_2(p+1)$, so that there exists a point on $E(\mathbb{F}_p)$ of exact order 2^{f-1} . Denote by T_0, T_{-1}, T_1 the three non-trivial 2-torsion points, with the indexes as above. Let $x_P \in \mathbb{F}_p$. Then

- $\begin{aligned} &-x_P \text{ lifts to a point } (x_P, y_P) = P \in E(\mathbb{F}_p) \text{ with } 2^{f-1} \mid \operatorname{ord}(P) \text{ if and only if} \\ &(x_P x(T_{-1})) \in \mathbb{F}_p^* \setminus \mathbb{F}_p^{*2} \text{ and } g(x_P) \in \mathbb{F}_p^{*2}, \\ &-x_P \text{ lifts to a point } (x_P, y_P) = P \in E^t(\mathbb{F}_p) \text{ with } 2^{f-1} \mid \operatorname{ord}(P) \text{ if and only if} \\ &(x_P x(T_1)) \in \mathbb{F}_p^{*2} \text{ and } g(x_P) \in \mathbb{F}_p^* \setminus \mathbb{F}_p^{*2}, \end{aligned}$

where

$$E^t/\mathbb{F}_p: by^2 = g(x)$$

denotes a quadratic twist of E over \mathbb{F}_p . (We remark that we can take b = -1because -1 is not a square over \mathbb{F}_p .)

Proof. For the first part, note that it is obvious that $P \in E(\mathbb{F}_p) \setminus E[2]$ if and only if $g(x_P) \in \mathbb{F}_p^{*2}$. Further, let $R \in E(\mathbb{F}_p)$ be any element of exact order 2^{f-1} . We can write $P = [u]R + [b]T_0 + S$ with S in $[2]E(\mathbb{F}_p)$. Now 2^{f-1} divides the order of P if and only if u is odd. The statement then follows from

$$e_{T,2}(T_{-1}, P) = e_{T,2}(T_{-1}, R)^u e_{T,2}(T_{-1}, T_0)^b = (-1)^u,$$

where $e_{T,2}(T_{-1}, R) = -1$ by combining Lemma D.1 and the non-degeneracy of the Tate pairing.

Next, write $\tau: E \to E^t$ for any twisting isomorphism leaving x fixed, and write T_{-1}^t, T_0^t, T_1^t for the non-trivial 2-torsion points on E^t , with indexes same as before. The second part of the lemma then follows from the first after noting that $\tau(T_1) = T_{-1}^t$, and that for any $T_{-1}^t \neq Q \in E^t(\mathbb{F}_p)$, the reduced Tate pairing $e_2(T_{-1}^t, Q)$ is given by the residue class of $\frac{x(Q) - x(T_{-1}^t)}{((x - x(T_{-1}^t))/(y/x)^2)(0_E)} = (x(Q) - x(Q))$ $x(T_{-1}^{t}))/b.$

Note that this gives a particularly effective algorithm for sampling a basis of the form we want.

Remark D.1 (Choosing a canonical representative for the Montgomery coeffi*cient*). Let E/\mathbb{F}_q be an elliptic curve. Recall (see *e.g.* [56, Example 5.12]) that if $T \in E[2](\mathbb{F}_q)$, E can be put in a Montgomery form $by^2 = x^3 + Ax^2 + x$ with T sent to (0,0) if and only if there exists a rational cyclic group of order $4 \ G \subset E[4]$

that contains T. Equivalently, T has trivial self Tate pairing. If that is the case, there are always exactly two different such subgroups, which give the two coefficients A, -A. As explained in Appendix B.4, since -1 is not a square over \mathbb{F}_p , then if b = -1, switching to -A allows us to always work with Montgomery models with b = 1.

There are always 6 possible Montgomery models over $\overline{\mathbb{F}}_p$ (two for each of the three 2-torsion points). However, in our situation of E/\mathbb{F}_p on the surface, only T_1 and T_{-1} give a rational Montgomery model. Indeed, T_0 induces the descending isogeny to the floor, which cannot be extended further to a rational 4-isogeny. Imposing the condition that b = 1, we only have two possible Montgomery coefficients for our elliptic curves.

In our implementation, we pick a deterministic one based on the lexicographic order. An alternative choice would be to (for instance) pick the choice that sends T_{-1} to (0,0). A further optimisation would be to represent the Montgomery curve not by the A coefficient, but by the x-coordinate α of T_0 (for instance), from which we recover A as $-\alpha - 1/\alpha$. With this convention, we would be able to pick up immediately the three points T_1, T_0, T_{-1} without a square root computation. For the class group action, since we first compute the theta coordinate of E_a , which encodes all the 2-torsion on E_a , we can easily output α_a rather than A_a . Explicitly, if (a:b) is a rational theta null point on E_a , then $\alpha_a = (b^2 - a^2)/(a^2 + b^2)$ is the x-coordinate of T_0 when E_a is put into the Montgomery form such that the 2-torsion theta point (-a:b) (which has trivial self pairing, hence is either T_1 or T_{-1}) is sent to (0:1).

D.2 Evaluating endomorphisms with denominator

In this section, we show how to evaluate endomorphisms with denominator, by using Tate-pairings instead of division points. We first give the general algorithm, before noting that our situation is particularly simple.

Algorithm 1 EvalEndomorphismWithDenominator(α, P)

Input: $\alpha \in \mathbb{Z}[\frac{1+\pi}{n}]$, where π denotes the *q*-power frobenius endomorphism, a point $P \in E(\mathbb{F}_q)$, where $\operatorname{End}_{\mathbb{F}_q}(E) = \mathbb{Z}[\frac{1+\pi}{n}]$

Output: $\alpha(P)$

1: Write $\alpha = \frac{a+b\pi}{n}$, with *n* of minimal denominator.

2: Compute a basis $\langle T_1, T_2 \rangle = E[n]$.

- 3: Compute the pairings $\mu_0 = e_{W,n}(T_1, T_2), \mu_1 = e_{T,n}(T_1, P), \mu_2 = e_{T,n}(T_2, -P).$
- 4: Compute r, s so that $\mu_0^r = \mu_2$ and $\mu_0^s = \mu_1$.
- 5: Set $T = [r]T_1 + [s]T_2$.
- 6: return $[b]T + [\frac{a+b}{n}]P$.

Proposition 19. Algorithm 1 is correct, and returns $\alpha(P)$ without using division points.

Proof. Since $\alpha - \frac{a+b}{n} = b\frac{\pi-1}{n}$, and $\frac{a+b}{n} \in \mathbb{Z}$, we see that evaluating $\alpha(P)$ reduces to some simple scalar multiplications and additions after evaluating $\frac{\pi-1}{n}(P)$.

Notice that $E[n] \subseteq E(\mathbb{F}_q)$, since $\pi - 1$ factors through multiplication by n. Thus, we can sample a basis $\langle T_1, T_2 \rangle = E[n]$ over \mathbb{F}_q . Let $\frac{\pi-1}{n}(P) = T$. We have that $[n]T = (\pi - 1)(T) = 0_E$, since $T \in E(\mathbb{F}_q) = \ker(\pi - 1)$, thus T can be written as $T = [r]T_1 + [s]T_2$ for some r, s. By the non-degeneracy of the Weil-pairing, we can recover r, s by computing the evaluations $e_{W,n}(T_1, T_2), e_{W,n}(T_1, T), e_{W,n}(T_2, T)$. Finally, by [56, Equation 15], we have that

$$e_{W,n}(T_i,T) = e_{T,n}(T_i,P),$$

which finishes the proof.

In the CSIDH/CSURF supersingular case, the maximal denominator n that can occur is n = 2, which is the situation we are in in Section 5.2. Further, we only ever need to evaluate the endomorphisms in multiples of our chosen basis points P, Q, where P, Q are points of order 2^{f-1} for $2^f || p + 1$

Notice that in this case, the algorithm becomes particularly easy; assuming we aim to compute $\alpha(R)$ for $\alpha = \frac{a+\pi b}{n}$, we are in one of two situations:

- If R = [m]P or R = [m]Q for some $m \equiv 0 \pmod{2}$, we see that the pairings $\mu_1 = \mu_2 = 1$, and hence $\alpha(R) = [\frac{a+b}{2}]R$.
- If R = [m]P or R = [m]Q for some $m \equiv 1 \pmod{2}$, $\frac{\pi-1}{2}(R)$ is instead given by the unique 2-torsion point T_i with $e_{T,2}(T_i, R) = 1$. But we already know that $e_{T,2}(T_{-1}, P) = e_{T,2}(T_1, Q) = -1$ by Lemma D.2, hence we see that a single Legendre test is enough to determine the correct T_i .

Example D.1. We of course have $\frac{\pi-1}{2}(T_1) = 0$, and by Lemma D.2, $\frac{\pi-1}{2}(T_0) = \frac{\pi-1}{2}(T_{-1}) = T_{-1}$ on E, and $\frac{\pi-1}{2}(T_0) = \frac{\pi-1}{2}(T_1) = T_1$ on E^t . If we sample P above T_1 , then if $e_{T,2}(T_0, P) = 1$, $\frac{\pi-1}{2}(P) = T_0$, whereas if $e_{T,2}(T_0, P) = -1$, $\frac{\pi-1}{2}(P) = T_1$.

Further, since translating by $T \in E[2]$ is still well defined on the Kummer line $E/\pm 1$, we see that evaluating our endomorphisms essentially only consists of a single scalar multiplication, and potentially a translation by T_i , which can all be done on the Kummer line.

Remark D.2. We can further refine our basis sampling to always output points such that $e_{T,2}(T_0, P) = e_{T,2}(T_0, Q) = 1$. By the discussion above, in this case we will always know that the potential translation by T_i is by $T_i = T_0$. Sampling such a basis is easy to do using Lemma D.2: we want that $x_P - x(T_0)$ is a square, *i.e.* $x_P = x(T_0) + u^2$, and since Q is on the twist, that $x_Q - x(T_0)$ is not a square, *i.e.* $x_Q = x(T_0) - u^2$. Then by the non degeneracy of the Tate pairing, we have $e_{T,2}(T_{-1}, P) = -1$ and $e_{T,2}(T_1, Q) = -1$, so by bilinearity we also have $e_{T,2}(T_1, P) = -1$ and $e_{T,2}(T_{-1}, Q) = -1$.

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D.3 Using the bottom of the volcano

We describe an alternative strategy to sample a basis on E by using the descending isogeny $\phi_{T_0}: E \to E_0$. Recall that T_0 is the unique 2-torsion point on E with non trivial self Tate pairing.

On E_0 , we are at the bottom of the 2-isogeny volcano, so the 2-Sylow is cyclic, generated by a rational point P_0 of order 2^f . The image of P_0 by $\hat{\phi}_{T_0}$ is then a point P of order 2^{f-1} on E. By construction of P, we have $e_{T,2}(T_0, P) = 1$, while $e_{T,2}(T_{-1}, P) = e_{T,2}(T_1, P) = -1$ because P is of full order on E.

Doing the same construction on E_0^t , we get a point $Q = \hat{\phi}_{T_0}(Q_0) \in E^t$, and (P, Q) is an appropriate basis (as in Remark D.2) to which to apply the evaluation algorithm of Appendix D.2.

We saw on Remark D.1 that there are 4 possible choice of Montgomery coefficient for E (or 2 if we restrict to b = 1). Also, because the 2-torsion is rational on E, then A + 2 being a square is equivalent to A - 2 being a square, in which case (1 : 1) and (-1 : 1) lift to a rational point of 4-torsion on $E : y^2 = x^3 + Ax^2 + x$. In the other case, then -A + 2 and -A - 2 are squares, and (1 : 1), (-1 : 1) lift to $E^t : y^2 = x^3 - Ax^2 + x$.

By contrast, on E_0 , there is only one rational point of 2-torsion T'_0 (which generates the kernel of $\hat{\phi}_{T_0}$), which has trivial self Tate pairing, so we can find a Montgomery form for E_0 with T'_0 sent to (0,0). Then E_0 is isomorphic to $by^2 = x^3 + A_0x^2 + x$, and replacing A_0 with $-A_0$ if necessary, we can assume b = 1. In particular, there is only one choice for A_0 to put E_0 in Montgomery form $E_0: y^2 = x^3 + A_0x^2 + x$. Furthermore, either $A_0 + 2$ is a square (so (1,1)lifts to E_0) and then $A_0 - 2$ is not a square (so (-1,1) does not lift), so $-A_0 + 2$ is a square and $-A_0 - 2$ being a square, $-A_0 + 2$ not a square, $-A_0 - 2$ a square.

We see that A_0 gives a convenient way to represent not only E_0 , but also E: we have $T'_0 = (0,0)$, $\phi_{T'_0}$ the canonical ascending isogeny $E_0 \to E$, whose dual ϕ_{T_0} has kernel generated by T_0 , and also that if $A_0 + 2$ is a square then the image of (1:1) by $\phi_{T'_0}$ gives T_1 , and the image of (-1:1) gives T_{-1} (and conversely if $A_0 + 2$ is not a square).

A last advantage is that we can efficiently sample $P_0 \in E_0(\mathbb{F}_p), Q_0 \in E^t(\mathbb{F}_p)$ of full order 2^f by adapting the entangled basis algorithm of [64]. We first sample $x(P_0)$ a non square in \mathbb{F}_p of the form $-A_0/(1-u^2)$ until we find that $x(P_0)$ lifts to a rational point P_0 on E_0 . Since the Tate pairing of T'_0 with P_0 is non trivial by construction, P_0 , multiplied by the cofactor, has exact order 2^f . Now set $x(Q_0) = -A_0 - x(P_0) = -u^2x(P_0)$. Then $x(Q_0)$ is a square, and $x(Q_0)^2 + A_0x(Q_0) + x(Q_0) = x(P_0)^2 + A_0x(P_0) + x(P_0) = y(P_0)^2/x(P_0)$ is not a square. So $x(Q_0)$ lifts to a point on the twist $-y^2 = x^3 + A_0x^2 + x$, with $y(Q_0) = uy(P_0)$. And the non reduced Tate pairing of T'_0 and Q_0 is given by the class of $-x(Q_0)$, which is not trivial. Hence Q_0 , multiplied by the cofactor, also has exact order 2^f on E^t . We can precompute a table of elements such that $-1/(1-u^2)$ is not a square. Then for the entangled basis generation, depending on whether A_0 is a square or not, we use the first or the second table. Remark D.3 (The horizontal isogenies of degree 2^{f-1}). On E, which is at the crater, the ideal (2) splits as $(2) = \mathfrak{p}_1\mathfrak{p}_{-1}$. We have $2^f \parallel \pi^2 - 1$, and $\mathfrak{p}_1^{f-1}\mathfrak{p}_{-1} \mid \pi - 1$, $\mathfrak{p}_{-1}^{f-1}\mathfrak{p}_1 \mid \pi + 1$.

Let (P_0, Q_0) be our basis sampled on E_0 as above, $P = \hat{\phi}_{T_0}(P_0), Q = \hat{\phi}_{T_0}(Q_0)$ the points of order 2^{f-1} on E. Then since T'_0 has non trivial pairing with P_0 , then by [56, Example 5.16], the isogeny generated by P_0 lands on the bottom of the volcano, hence $\langle P \rangle$ is not the kernel of the isogeny associated to \mathfrak{p}_1^{f-1} , as can also be seen from the fact that it has non trivial pairing with T_1 . However, if $P' = P + T_{-1}$ or $P' = P + T_0$, then $e_{T,2}(T_1, P') = 1$ by Lemma D.1, and so $\langle P' \rangle$ is the kernel of \mathfrak{p}_1^{f-1} . A similar argument shows that $E[\mathfrak{p}_{-1}^{f-1}] = \langle Q + T_0 \rangle$.

Remark D.4 (The 2-isogenies between E and E_0). As illustrated above, it is convenient to use the 2-isogenies ϕ_{T_0} and $\phi_{T'_0}$ to move between E and E_0 . We also would like to have E in Montgomery form, and E_0 too for the entangled basis generation. To put a curve E in Montgomery form, we need the x-coordinate of a point of 4-torsion $T' \in E$ such that $x(T') \in \mathbb{F}_p$. In particular, if we start with E_0 in Montgomery form, and take the quotient by T'_0 , we easily find E in Legendre form (see for instance [58, Example B.1]), but finding a Montgomery form for E requires either a square root or to push the 8-torsion point $2^{f-3}P_0$ (or $2^{f-3}Q_0$) to E via $\phi_{T'_0}$.

On the other hand, if we start from E in Montgomery form and we are given $\alpha = x(T_0)$, then we can directly find E_0 in Montgomery form using the formulas from [51, Proposition 2]. Hence it may be useful to use α to represent E (or even use the level 2 theta null point), as already suggested in Remark D.1, to save a square root.