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¡Abstract here;

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Chapter 1

Introduction

- 1.1 Motivation
- 1.1.1 Literature Survey
- 1.2 Contributions
- 1.3 Structure

1.3.1 Layout

Over the course of the past decade, elliptic curve cryptography (ECC) has proven itself a mainstay in the wide world of applied cryptology. While isogeny based cryptography does build itself up from the same underlying field of mathematics as ECC, it simultaneously draws from a slightly more complicated space of algebraic notions. Much of this chapter will be dedicated to illuminating these notions in a manner that should be digestable for those without serious background in algebraic geometry, or abstract algebra in general.

1.3.2 Notation

Chapter 2

Technical Background

This chapter will cover the following preliminary topics: cryptographic primitives, isogenies and their relevant properties, supersingular isogeny Diffie-Hellman (SIDH), the Fiat-Shamir construction for digital signatures (and its quantum-safe adaptation), the current landscape of isogeny based signature schemes, and finally the C implementations of isogeny based protocols with which we are concerned.

In the first section of this chapter we will take some time to introduce a few ideas from modern cryptography. We will cover key exchange, identification schemes, and signature schemes - all at as high of an abstraction level as possible. Readers familiar with these topics can skip this section without harm.

Our discussion of isogenies will begin with some basic coverage of the underlying algebra. We will provide the material necessary for the remaining sections as we build up in the level of abstraction; working our way through groups, finite fields, elliptic curves, and finally isogenies and their properties.

Once we have presented the necessary algebra, we will illustrate the specifics of the supersingular isogeny Diffie-Hellman key-exchange protocol. We will spend most of this time dedicated to a modular deconstruction of the protocol, looking at the high-level procedures and algorithms which will be necessary for understanding in detail the signature protocol to come. This subsection will end with a briefing and analysis of the closely related zero-knowledge proof of identity (ZKPoI) isogeny protocol proposed in the original De Feo et al. paper [FJP12], as it is the foundation for the isogeny based signature scheme presented by Yoo et al [YAJ⁺12].

In section 2.3 we will discuss the Fiat-Shamir transformation [Kat10]; a technique which, given a secure interactive identification scheme, creates a secure digital signature scheme. We will also look at the quantum-secure adaptation published by Unruh [Unr14] for applying a non-quantum-resistant transform to a quantum-resistant primitive would be rather frivolous.

Section 2.4 will be dedicated to covering current isogeny-based signature schemes - the topic about which this dissertation is mainly concerned. We will discuss the signature scheme of Yoo et al., which is a near direct application of Unruh's work to the SIDH zero-knowledge proof of identity.

Finally, the last section of this chapter will introduce the SIDH C library released by Microsoft Research, on top of which the core contributions of this thesis are implemented. We will also look at the implementation of the to-be-discussed signature scheme, which is a proof-of-concept built on top of the Microsoft API.

2.1 Cryptographic Primitives

Cryptographic primitives can be thought of as the basic building blocks used in the design of cryptographically secure applications and protocols. The idea of which being that if individual primitives are provably (or believeably) secure, we can be more confident in the security of the application as a whole.¹

To quickly recap some basic information security, there are serveral different security properties a cryptographic primitive may aim to offer:

- Confidentiality: The notion that the information in question is kept private from unauthorized individuals.
- *Integrity*: The notion that the information in question has not been altered by unauthorized individuals.
- Availability: The notion that the information in question is available to authorized individuals when requested.
- Authenticity: The notion that the source of the information in question is verified.
- Non-repudiation: The notion that the source of the information in question **cannot** deny having originally provided the information.

The security of a particular cryptographic primitive is measured by two components. The first, referred to often as a "security guarantee", measures what conditions constitute a successful attack on the primitive. The second, known as the "threat model", makes assumptions about the computational powers that the adversary holds. The best practice in forming security proofs is to aim for security with respect to the most easily broken security guarantee and the most challenging possible threat model. The combination of a security guarantee and threat model is known as a security goal.

Each of the primitives to come are designed to offer some utility in the communication between a given pair of entities. We will refer to these entities as Alice and Bob. The schemes we are concerned with in this dissertation are strictly *public key* (also known as *asymmetric key*) schemes. In public key primitives, each user possesses a *public* key (visible to every user in the network) as well as a *private* key, which only they have access to.

The first class of primitives we will discuss, *key exchange* protocols, provide a means by which Alice and Bob can come to the agreement of some secret value. The goal of a key exchange protocol is for Alice and Bob, communicating over some open, insecure channel, to reach mutual agreement of the secret value while also ensuring the *confidentiality* of that value. The secret value is referred to as a *secret* or *shared* key and is intended for use in other cryptographic primitives.

Identification schemes are a class of primitives that aim to ensure *authenticity* of a given entity. If Alice is communicating with Bob and she wants to verify that Bob is who he claims to be, the two can utilize a secure identification scheme. After identification protocols we will look at signature schemes, which are somewhat of an extension of the former. Signature schemes aim to provide *authenticity* on every message sent from Bob

¹This is not to say that software which implements provably secure primitives is guaranteed to be secure. In security, it should be expected that the weakest link in the system is the first to be exploited, and these weak links often lie in careless implementation details.

to Alice, as well as non-repudiability and integrity of those messages.

Random Oracle Model. Before continuing with our discussion of primitives, it is worth discussion a framework in cryptography known as the random oracle model. A "random oracle" is a theoretical black box which, for every unique input, responds with a truly random output. That is, if a query is made to a random oracle h with input x (written h(x)) multiple times, h with respond with the same (random) output every time.

For certain constructions to be proven secure, it is sometimes necessary or helpful to assume the existence of random oracles. While this assumption may seem greviously optimistic, *hash functions* are a widely diployed family of functions which are believed to approach the nature of random oracles to some degree. Much of the security of modern cryptography depends on the security of such hash functions.

2.1.1 Key Exchange

A key exchange protocol, which we will denote as Π_{kex} , can be represented in some contexts by a pair of polynomial time algorithms **KeyGen** and **SecAgr**: $\Pi_{kex} = (\mathbf{KeyGen}, \mathbf{SecAgr})$. Alice and Bob will each run both of these procedures. The first they will run on the same input, 1^{λ} , a bit string of λ 1's. The second, short for "secret agreement", they will run on both their outputs of **KeyGen** and their peers.

Execution of Π_{kex} between Alice and Bob involves the following:

- (i) Alice and Bob run $\mathbf{KeyGen}(1^{\lambda})$: A probabilistic algorithm with input 1^{λ} and output (sk, pk). Typically pk is the image of f(sk), where f is some *one-way* function. We will denote the outputs of \mathbf{KeyGen} for Alice and Bob as (sk_{Alice}, pk_{Alice}) and (sk_{Bob}, pk_{Bob}) respectively.
- (ii) Alice and Bob exchange (over an insecure channel) their public keys pk_{Alice} and pk_{Bob} .
- (iii) Alice runs $\mathbf{SecAgr}(sk_{Alice}, pk_{Bob})$: A deterministic algorithm with input sk_{Alice} and pk_{Bob} and output $k_{Alice} \in \{0, 1\}^{\lambda}$. Bob runs $\mathbf{SecAgr}(sk_{Bob}, pk_{Alice})$ to obtain $k_{Bob} \in \{0, 1\}^{\lambda}$.

 Π_{kex} is said to uphold *correctness* if $k_{Alice} = k_{Bob}$ for all honestly derived keypairs (sk_{Alice}, pk_{Alice}) and (sk_{Bob}, pk_{Bob}) . Because we deal only with correct Π_{kex} , we refer to the output of Π_{kex} as simply k. Figure 2.1 illustrates an execution of the Diffie-Hellman key exchange protocol which relies on the difficulty of the *discrete logarithm* problem for its one-way function f.

The security goal typical of a key exchange protocol is that an adversary with access to the session transcript (threat) cannot discern the resulting shared secret key from a randomly generated value (security guarantee).

2.1.2 Interactive Identification Schemes

Imagine Alice wishes to confirm the identity of Bob. The motivation for interactive identification schemes is to provide Bob with some mechanism for proving to Alice (or any other party) that he has knowledge of some secret which **only** Bob could possess.

Public parameter:



Figure 2.1: Alice and Bob's execution of Diffie-Hellman key exchange.

The goal, of course, being to accomplish this without openly revealing the secret, so that it can continue to be used as an identifier for Bob.

An identification scheme (or otherwise "proof of identity" protocol) Π_{id} is composed of by the tuple of polynomial-time algorithms (**KeyGen**, **Commit**, **Prove**, **Verify**) and some set ω . Π_{id} is an interactive protocol requiring two parties. The *prover* (Bob, for example) executes **KeyGen**, **Commit**, and **Prove**. The *verifier* (Alice, in our example) executes **Verify** following Bob's actions.

Execution of Π_{id} between Alice and Bob proceeds as follows:

- (i) Bob runs **KeyGen**(1^{λ}): A probabilistic algorithm with input 1^{λ} and outputs Bob's keypair (sk, pk). Bob sends his public key pk to Alice.
- (ii) Bob runs **Commit()**: a probabilistic algorithm with output *com*. *com* is referred to as a "commitment". Bob sends *com* to Alice.
- (iii) Alice sends a randomly generated "challenge" value $ch \in \omega$. Alice sends ch to Bob.
- (iv) Bob runs Prove(sk, com, ch): A deterministic algorithm with input sk and ch, and output resp. resp is the "response" to Alice's challenge.
- (v) Alice runs Verify(pk, com, ch, resp): A deterministic algorithm with input pk, com, ch, and resp, and output $b \in 0, 1$. Bob has successfully proven his identity to Alice if b = 1.

If Alice accepts Bob's response, and b=1, then we refer to the tuple (com, ch, resp) as an accepting transcript. This general construction for identification protocols is illustrated in Figure 2.2, where the prover is referred to as \mathcal{P} and \mathcal{V} denotes the verifier.

In terms of security, it is common to show that an identification scheme is secure against *impersonation* under a *passive attack*. Proving such security implies that an adversary who eavesdrops on arbitrarily many executions of Π_{id} between a verifier \mathcal{V} and a prover \mathcal{P} cannot successfully impersonate \mathcal{V} .

We may at times speak of *canonical* identification schemes. An identification scheme Π occurring between a prover \mathcal{P} and a verifier \mathcal{V} is labelled canonical if it satisfies all of the following constraints:

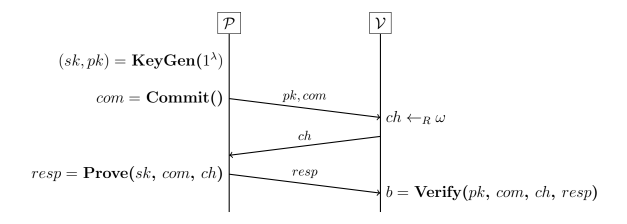


Figure 2.2: A general interactive identification scheme with prover \mathcal{P} and verifier \mathcal{V} .

- Π consists of an initial message (or "commitment") com sent by \mathcal{P} , a challenge ch sent by \mathcal{V} , and a final response resp sent by \mathcal{P} .
- ch is chosen uniformly at random from some set ω .
- com is generated by some probabilistic function **R** taking \mathcal{P} 's secret key as input. For any secret key sk and fixed string $c\bar{o}m$, the probability that $\mathbf{R}(sk) = c\bar{o}m$ is negligible.

These constraints gurantee that Π will have two important features. First, that any third party given the transcript and prover's public key can efficiently determine whether the verifier will accept. Second, that the probability that *comm* repeats in polynomially many executions of Π is negligible.

Lastly, it should be mentioned that there exist variations upon this type of primitive wherein Alice is not required to send Bob a specific challenge value. These are known as *non-interactive* identification schemes, or non-interactive proofs of identity (NIPoI). These non-interactive approaches to solving the problem of identity and *authentication* further bridge the gap between identification protocols and signature schemes.

2.1.3 Signature Schemes

We define a signature scheme as the tuple of algorithms $\Pi_{sig} = (\mathbf{KeyGen}, \mathbf{Sign}, \mathbf{Verify})$. Some execution of Π_{sig} between Alice and Bob for a particular message m sent from Bob to Alice involves the following... First, before any message is to be signed, the Bob must run the following:

• Bob runs **KeyGen**(1^{λ}): A probabilistic algorithm with input 1^{λ} and output (sk, pk).

Then, for every message m Bob wishes to authenticate and send to Alice:

- (i) Bob sends his public key pk to Alice over an authenticated channel if he has not yet done so.
- (ii) Bob runs $\mathbf{Sign}(sk, m)$: A probabilistic algorithm with input sk (Bob's secret key) and m (the message Bob wishes to authorize) and output σ , known as a *signature*.
- (iii) Bob sends m and σ to Alice.

(iv) Alice runs $\mathbf{Verify}(pk, m, \sigma)$: A deterministic algorithm with input pk (Bob's public key), m, and σ and output $b \in \{0, 1\}$. Alice has confidence in the *integrity* and origin authenticity of m if b = 1.

As previously alluded to, it is worth noting that signature protocols and identification schemes are closely related. In essence, they are rather similar; but with two main differences. The first is rather comparable to the aforementioned difference between interactive identification schemes and non-interactive identification schemes. The second arises as a result of aiming to authenticate Bob on any particular message m. To achieve this, the signature scheme needs to be run every time Bob wishes to send a message to Alice. The details of this comparison are intentionally left vague, as it will from a topic of close inspection in section 2.4.

The strongest security goal for a signature scheme Π_{sig} is expressed as existential unforgeability under an adaptive chosen-message attack. If this goal is provably satisfied, an adversary with the ability to sign arbitrary messages will not be able to forge any conceivable and valid signature.

2.2 Algebraic Geometry & Isogenies

Groups & Varieties. A group is a 2-tuple composed of a set of elements and a corresponding group operation (also referred to as the group law). Given some group defined by the set G and the operation \cdot (written as (G, \cdot)) it is typical to refer to the group simply as G. If \cdot is equivalent to some rational mapping $^2f_G: G \to G$, then the group (G, \cdot) is said to form an algebraic variety. A group which is also an algebraic variety is referred to as an algebraic group.

G is said to be an *abelian* group if, in addition to the four traditional group axioms (closure, associativity, existence of an identity, existence of an inverse), G satisfies the condition of commutativity. More formally, for some group G with group operation \cdot , we say G is an abelian group iff $x \cdot y = y \cdot x \ \forall x, y \in G$. An algebraic group which is also abelian is referred to as an **abelian variety**.

Definition 1 (Abelian Variety). for some algebraic group G with operation \cdot , we say G is an abelian variety iff $x \cdot y = y \cdot x \ \forall x, y \in G$.

For some group (G, \cdot) , some $x, y \in G$, and some rational mapping $f_G : G \to G$, let the following sequence of implications denote the classification of (G, \cdot) :

group
$$\xrightarrow{x \cdot y = f_G(x,y)}$$
 algebraic group $\xrightarrow{x \cdot y = y \cdot x}$ abelian variety

Morphisms. Let us again take for example some group (G, \cdot) . Let's also define some set $S_{(G,\cdot)}$ which contains every tuple (x, y, z) for group elements x, y, z which satisfy $x \cdot y = z$.

$$S_{(G,\cdot)} = \{x, y, z \in G | x \cdot y = z\}$$

Take also for example a second group (H,*) and some map $\phi: G \to H$. ϕ is said to be structure preserving if the following implication holds:

$$(x,y,z) \in S_{(G,\cdot)} \Rightarrow (\phi(x),\phi(y),\phi(z)) \in S_{(H,*)}$$

²A rational map is a mapping between two groups which is defined by a polynomial function with rational coefficients.

A **morphism** is simply the most general notion of a structure preserving map. More specifically, in the domain of algebraic geometry, we will be dealing with the notion of a **group homomorphism**, defined as follows:

Definition 2 (Group Homomorphism). For two groups G and H with respective group operations \cdot and *, a group homomorphism is a structure preserving map $h: G \to H$ such that $\forall u, v \in G$ the following holds:

$$h(u \cdot v) = h(u) * h(v)$$

From this simple definition, two more properties of homomorphisms are easily deducible. Namely, for some homomorphism $h: G \to H$, the following properties hold:

- h maps the identity element of G onto the identity element of H, and
- $\bullet \ h(u^{-1}) = h(u)^{-1}, \forall u \in G$

Recall that for some morphism (or function) $h: G \to H$, we refer to G as the domain and H as the codomain.

Furthermore, an *endomorphism* is a special type of morphism in which the domain and the codomain are the same groups. We denote the set of enomorphisms defineable over some group G as $\operatorname{End}(G)$. The *kernel* of a particular homomorphism $h:G\to H$ is the set of elements in G that, when applied to h, map to the identity element of H. We write this set as $\ker(h)$, and it is much analogous to the familiar concept from linear algebra, wherein the kernel denotes the set of elements mapped to the zero vector by some linear map.

2.2.1 Fields & Field Extensions

A field is a mathematical structure which, while being similar to a group, demands additional properties. Fields are defined by some set F and two operations: addition and multiplication. In order for some tuple $(F, +, \cdot)$ to constitute a field, it must satisfy an assortment of axioms:

Addition axioms:

- (closure) If $x \in F$ and $y \in F$, then $x + y \in F$.
- \bullet + is commutative.
- + is associative.
- F contains an element 0 such that $\forall x \in F$ we have 0 + x = x.
- $\forall x \in F$ there is a corresponding element $-x \in F$ such that x + (-x) = 0.

Multiplication axioms:

- (closure) If $x \in F$ and $y \in F$, then $x \cdot y \in F$.
- · is commutative.

- is associative.
- F contains an element $1 \neq 0$ such that $\forall x \in F$ we have $x \cdot 1 = x$.
- $\forall x \neq 0 \in F$ there is a corresponding element $x^{-1} \in F$ such that $x \cdot (x^{-1}) = 1$.

Additionally, a field $(F, +, \cdot)$ must uphold the distributive law, namely:

$$x \cdot (y+z) = x \cdot y + x \cdot y \text{ holds } \forall x, y, z \in F$$

While these axioms are known to be satisfied by the sets \mathbb{Q} , \mathbb{R} , and \mathbb{C} with typically defined + and \cdot , our focus will be on a particular class of field known as a *finite field*. Finite fields, as the name suggests, are fields in which the set F contains finitely many elements - we refer to the number of elements in F as the *order* of the field.

Let us take some prime number p. We can construct a finite field by taking F as the set of numbers $\{0, 1, ... p - 1\}$ and defining + and \cdot as addition and multiplication *modulo* p. Finite fields defined in this fashion are denoted as \mathbb{F}_p , and have order p.

$$\forall x, y \in \mathbb{F}_p, \ x + y = (x + b) \mod p, \text{ and }$$

 $\forall x, y \in \mathbb{F}_p, \ x \cdot y = (x \cdot b) \mod p$

For any given field K there exists a number q such that, for every $x \in K$, adding x to itself q times results in the additive identity 0. This number is referred to as the characteristic of K, for which we write $\operatorname{char}(K)$. Finite fields are the only type of field for which $\operatorname{char}(K) > 0$. Furthermore, if the field in question is finite and has prime order, then the order and the characteristic are equivalent.

A particular field K' is called an extension field of some other field K if $K \subseteq K'$. The complex numbers \mathbb{C} , for example, are an extension field of \mathbb{R} . A given field K is algebraically closed if there exists a root for every non-constant polynomial defined over K. If K itself is not algebraically closed, we denote the extension of K that is by \overline{K} .

An algebraic group G_a is defined over a field K if each element $e \in G_a$ is also an element of the field K, and the corresponding f_{G_a} is defined over K. To show that a particular algebraic group G_a is defined over some field K we will henceforth denote the group/field pairing as $G_a(K)$. Naturally, in the case where our field is a finite field of order p, we write $G_a(\mathbb{F}_p)$.

These algebraic structures are all important for building up to the concept of an *isogeny*. The lowest-level object we will be concerned with when discussing the forthcoming isogeny-based protocols will typically be elements of abelian varieties. The lowest-level structure in the SIDH C codebase is a finite field element.

Montgomery Arithmetic. We will now briefly discuss a technique for efficiently performing modular arithmetic. This method is widely deployed in cryptosystems centered around finite fields, and is abundantly used in the SIDH_C library that we will shortly be examining.

In 1985, Peter Montgomery introduced a method for efficiently computing the modular multiplication of two elements a and b. The technique begins with the construction of some constant R, whose value depends solely on the modulus N and the underlying computer architecture.³

³The specifics of how R is constructed are beyond the scope of this dissertation; if they feel so inclined, the reader should refer to [Mon85]

With the retrieval of R, $aR \mod N$ and $bR \mod N$ are constructed and referred to as the Montgomery representation of a and b respectively. Montgomery multiplication outlines an algorithm for computing $abR \mod N$ (the Montgomery product of a and b), from which $ab \mod N$ can be recovered through conversion back to standard representation. Once in Montgomery representation, other arithmetic can be performed (including field element inversions) in order to leverage the performance improvement offered by Montgomery modular multiplication – converting back to regular representation when necessary.

Applying Montgomery multiplication has the added benefit of decreasing the amount of field element inversions that need to be computed. Because of this, the technique is particularly relevant to this dissertation. We continue this discussion in section 3.2.

2.2.2 Elliptic Curves

An elliptic curve is an algebraic curve defined over some field K, the most general representation of which is given by

$$y^2 + a_1 xy + a_3 y = x^3 + a_2 x^2 + a_4 x + a_6.$$

This representation encapsulates elliptic curves defined over any field. If, however, we are discussing curves defined specifically over a field K such that $\operatorname{char}(K) > 3$ (see [Sil]), then the more compact form $y^2 = x^3 + ax + b$ can be applied (see Figure 2.3 for a geometric visualization). In this dissertation we will default to this second representation, as the schemes with which we are concerned will always be defined over \mathbb{F}_p for some large prime p.

We can define a group structure over the points of a given elliptic curve (or any other smooth cubic curve). If we wish to define a group in accordance to a particular curve, we do so with the following notation:

$$E: y^2 = x^3 + ax + b$$

Wherein E denotes the group in question, the elements of which are all the points (solutions) of the curve. Throughout much of this section, the words point and element can be used interchangeably.

The Group Law. The group operation we define for E, denoted +, is better understood geometrically than algebraically. Consider the following.

Given two elements P and Q of some arbitrary elliptic curve group E, we define + geometrically as follows: drawing the line L through points P and Q, we follow L to its third intersection on the curve (which is guaranteed to exist), which we will denote as $R = (x_R, y_R)$. We then set P + Q = -R, where -R is the reflection of R over the x-axis: $(x_R, -y_R)$. This descriptive definition of + is suitable for all situations except for when L is tangent to E or when E is parallel to the y-axis. These cases will be covered in a short moment. See Figure 2.3 for an illustrated representation of this process.

The group operation + is referred to as *pointwise addition*. In order for (E, +) to properly form a group under pointwise addition, it must satisfy the four group axioms:

• Closure: Because elliptic curves are polynomials of degree of 3, we know any given line passing through two points P and Q of E will pass through a third point R. The exceptions here are twofold. First, when P = Q and thus our line is tangent to

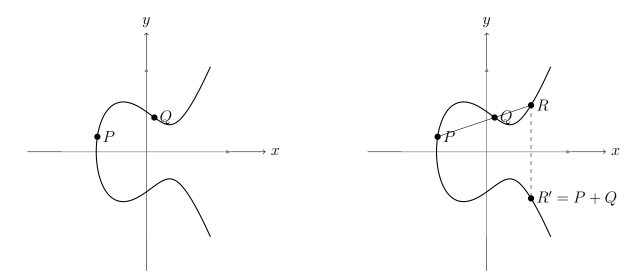


Figure 2.3: + acting over points P and Q of $y^2 = x^3 - 2x + 2$.

E, and second, when Q = -P and our line is parallel with the y-axis. We resolve the first case nicely by defining P + P by means of taking L to be the line tangent to E at point P. In the second case, P + (-P), by group axiom, should yield the identity element of the group. We will define this element and resolve this issue below.

- Identity: The identity element of elliptic curve groups, denoted as \mathcal{O} , is a specially defined point satisfying $P + \mathcal{O} = \mathcal{O} + P = P$, $\forall P \in E$. Because of the inclusion of this special element, we have that #(E(K)) is equal to 1 + the number geometric points on E defined over K. This of course is only a noteworthy claim when K is a finite field (otherwise there are already infinitely many elements in E).
- Associativity: For all points P, Q, and R in E, it must be the case ((P+Q)+R=P+(Q+R)) holds. It is rather easy to see visually why this is true for geometrically defined points in E (see Figure 2.4). Additionally, we can trivially show that this holds when any combination of P, Q, and R are \mathcal{O} by applying the axiom of the identity.
- Inverse: Due to the x-symmetry of elliptic curves, every point $P = (x_P, y_P)$ of E has an associated point $-P = (x_P, -y_P)$. If we apply + to P and -P, L assumes the line parallel to the y-axis at $x = x_P$. As discussed above, in this case there is no third intersection of L on E. In light of this, \mathcal{O} can be thought of as a point residing infinitely far in both the positive and negative directions of the y-axis. \mathcal{O} is equivalently referred to as the point at infinity (see Figure 2.4).

Of course, there are relatively simple formulas for algebraically defining point-wise addition and inverse computation. We have opted to describe these operations geometrically simply for ease of communication.

⁴One might suspect that the inclusion of this (apparently) non-algebraic element \mathcal{O} suggests that + is not a rational-map. The operator + can be shown to be a rational-mapping if we define our elliptic curve groups in three-dimensional projective space.



Figure 2.4: associativity illustrated on $y^2 = x^3 - 3x$ (left & center) and $P + (-P) = \mathcal{O}$ illustrated for $y^2 = x^3 + x + 1$ (right).

Additionally, we shorthand P + P + ... + P as nP, analogous to scalar multiplication.

Consequently, because groups defined over elliptic curves in this fashion are commutitive, they also constitute abelian varieties.

When referring to curves as abelian varieties defined over a field, we will write them as $E_{\alpha}(K)$, for some curve E_{α} and some field K. If we are only concerned with the geometric properties of the curve, or curves as distinct elements of some group structure, it will suffice to write E_{α} . Moving forward from here, we will assume all general curves discussed are capable of definition over some finite field \mathbb{F}_p .

The r-torsion group of E is the set of all points $P \in E(\overline{\mathbb{F}}_q)$ such that $rP = \mathcal{O}$. We denote the r-torsion group of some curve as E[r].

Supersingular Curves. An elliptic curve can be either ordinary or supersingular. There are several equivalent ways of defining supersingular curves (and thus the distinction between them and ordinary curves) in a general setting, but each of these goes well beyond our scope. In the context of curves defined over finite fiels, however, the following succinct definition holds:

Definition 3 (Supersingular Curve). Let E be an elliptic curve defined over the finite field \mathbb{F}_p . E is said to be supersingular if $\#(E(\mathbb{F}_p)) = p + 1$.

For the remainder of this paper, unless otherwise noted, all elliptic curves in discussion will supersingular.

Projective Space. While elliptic curves are naturally defined in two-dimensional affine space, there are many benefits to expressing them through three-dimension projective coordinates. First and foremost, expressing curves in projective space allows us to reason geometrically about \mathcal{O} . This is done by defining a curve E such that it resides in some two-dimension subspace of 3-space, the point \mathcal{O} then resides at some point in 3-space outside of the residing plane of E.

^aReaders are welcomed to investigate [Cos] for further details.

Representing a curve in 3-space requires some substitution of x and y coordinates, a typical forma for achieving this is the following:

$$x = X/Z$$
 $y = Y/Z$ $Z = 1$

Such a representation of elliptic curve points offers more computationally efficient arithmetic over points. This is conceptually similar to the previously mentioned Montgomery arithmetic regarding finite field elements. Other substitutions offer different computational advantages, but the implementations we will discuss make use of this typical approach.

2.2.3 Isogenies & Their Properties

Definition 4 (Isogeny). Let G and H be algebraic groups. An <u>isogeny</u> is a morphism $h: G \to H$ possessing a finite kernel.

In the case of the above definition where G and H are abelian varieties (such as elliptic curves,) the isogeny h is homomorphic between G and H. Because of this, isogenies over elliptic curves (and other abelian varieties) inherit certain characteristics.

For an isogeny $h: E_1 \to E_2$ defined over elliptic curves E_1 and E_2 , the following holds:

- $h(\mathcal{O}) = \mathcal{O}$, and
- $h(u^{-1}) = h(u)^{-1}, \forall u \in G$

If there exists some isogeny ϕ between curves E_1 and E_2 then E_1 and E_2 are said to be *isogenous*. All supersingular curves are isogenous only to other supersingular curves. The equivalent statement holds for ordinary curves. With this in mind, we can concieve a sort of graph structure connecting all isogenous curves, these graphs pertaining to either the supersingular or ordinary variety of curves.

We write $\operatorname{End}(E)$ to denote the ring formed by all the isogenies acting over E which are also endomorphisms. Note that m-repeated pointwise addition of a point with itself can equivalently be modelled by an endomorphism, we denote the application of such an endomorphism to a point P as [m]P, such that $[m]: E \to E$ and [m]P = mP.

An important facet of isogenies is that they can be uniquely identified by their kernel. If S is the group of points denoting the kernel of some isogeny ϕ with domain E, we write $\phi: E \to E/S$. Because the subgroup S sufficiently identifies ϕ , any given generator of S equivalently identifies ϕ . Therefore, if R generates the subgroup S we can write $\phi: E \to E/\langle R \rangle$. Moreover, we will have a specific interest in isogenies with kernels defined by some $torsion\ subgroup$.

Lemma 1 (Uniquely identifying isogenies). Let E be an elliptic curve and let Φ be a finite subgroup of E. There are a unique elliptic curve E' and a separable isogeny $\phi: E \to E'$ satisfying $ker(\phi) = \Phi$.

2.3 Supersingular Isogeny Diffie-Hellman

This section will aim to accomplish two things. First, we will briefly explain the isogeny-level & key-exchange-level procedures of the SIDH protocol. Second, we will illuminate

how these procedures map onto Microsoft Research's C implementation of SIDH. In this regard, this section can be considered an attempt to meld two domains of SIDH functions & procedures, in hopes of easing the navigation from the SIDH protocol to Microsoft's C implementation, and vice versa.

The original work of De Feo, Jao, and Plut [FJP12] outlines three different isogeny-based cryptographic primitives: Diffie-Hellman-esque key exchange, public key encryption, and the aforementioned zero-knowledge proof of identity (ZKPoI). Because all three of these protocols require the same initialization and public parameters, we will begin by covering these parameters in detail. Immediately after, we will analyze the key exchange at a relatively high level. Our goal of this section is to explain in detail the algorithmic and cryptographic aspects of the ZKPoI scheme, as this forms the conceptual basis for the signature scheme we will be investigating. We begin with the key exchange protocol because its sub-routines are integral to the Yoo et al. signature implementation.

For the discussion that follows, we will assume every instance of an SIDH protocol occurs between two parties, A and B (eg. Alice & Bob,) for which we will colorize information particular to A in red and B in blue. This will include private keys & public keys as well as the variables and constants used in their generation.

Public Parameters. As the name suggests, SIDH protocols work over supersingular curves (with no singular points). Let $\mathbb{F}_q = \mathbb{F}_{p^2}$ be the finite field over which our curves are defined, \mathbb{F}_{p^2} denoting the quadratic extension field of \mathbb{F}_p . p is a prime defined as follows:

$$p = \ell_A^{e_A} \ell_B^{e_B} \cdot f \pm 1$$

Wherein ℓ_A and ℓ_B are small primes (typically 2 & 3, respectively) and f is a cofactor ensuring the primality of p. We then define globally a supersingular curve E_0 defined over \mathbb{F}_q with cardinality $(\ell_A^{e_A}\ell_B^{e_B}f)^2$. Consequently, the torsion group $E_0[\ell_A^{e_A}]$ is \mathbb{F}_q -rational and has $\ell_A^{e_A-1}(\ell_A+1)$ cyclic subgroups of order $\ell_A^{e_A}$, with the analogous statement being true for $E_0[\ell_B^{e_B}]$. Additionally, we include in the public parameters the bases $\{P_A, Q_A\}$ and $\{P_B, Q_B\}$, generating $E[\ell_A^{e_A}]$ and $E[\ell_B^{e_B}]$ respectively.

This brings our set of global parameters, G, to the following:

$$G = \{p, E_0, \ell_A, \ell_B, e_A, e_B, \{P_A, Q_A\}, \{P_B, Q_B\}\}$$

2.3.1 SIDH Key Exchange

This subsection will illustrate an SIDH key exchange run between party members Alice and Bob. The general idea of the protocol is summerized in the diagram below. In the scheme, **private keys** take the form of isogenies defined with domain E, and **public keys** are the associated co-domain curve of said isogenies.[FJP12] For the entirety of this section we will denote isogenies by their function symbols, but when we come section 2.6 to will show how we can efficiently represent isogenies in a computational environment.

$$E_{0} \xrightarrow{\phi_{A}} E_{0}/\langle A \rangle$$

$$\phi_{B} \downarrow \qquad \qquad \downarrow \phi'_{B}$$

$$E_{0}/\langle B \rangle \xrightarrow{\phi'_{A}} E_{0}/\langle A, B \rangle$$

The premise of the protocol is that both parties each generate a random point (A or B in the diagram,) which, according to proposition 1, identifies some distinct isogeny $\phi_A : E_0 \to E/\langle A \rangle$ (or equivalent for B). Alice and Bob then exchange codomain curves and compute

$$\phi_{\mathbf{A}}(E_0/\langle B \rangle)
\text{or}
\phi_{\mathbf{B}}(E_0/\langle \mathbf{A} \rangle).$$

From these isogenies, Alice and Bob arrive at their shared secret agreement: the mutual codomain curve of $\phi_A(E_0/\langle B \rangle)$ and $\phi_B(E_0/\langle A \rangle)$, denoted E_{AB} .

Below we've outlined the SIDH key exchange protocol $\Pi_{\text{SIDH}} = (\text{KeyGen}, \text{SecAgr})$ in a descriptive manner. We do not provide algorithmic definitions for all of these procedures, but algorithmic details for some of these are covered partly in Sections 2.6 and 3.2. C code for functions that are not covered in these Sections but are noneless relevant can be found in Appendix A.

KeyGen(λ): Alice chooses two random numbers $m_A, n_A \in \mathbb{Z}/\ell_A^{e_A}\mathbb{Z}$ such that $(\ell_A \nmid m_A) \vee (\ell_A \nmid n_A)$. Alice then computes the isogeny $\phi_A : E_0 \to E_A$ where $E_A = E_0/\langle [m_A]P_A, [n_A]Q_A \rangle$ (equivalently, $\ker(\phi_A) = \langle [m_A]P_A, [n_A]Q_A \rangle$). Bob does the same for random elements $m_B, n_B \in \mathbb{Z}/\ell_B^{e_B}\mathbb{Z}$.

Alice then applies her isogeny to the points which Bob will use in the creation of of his isogeny: $(\phi_A(P_B), \phi_A(Q_B))$. Bob performs the analogous operation. This leaves us with the following private and public keys for Alice and Bob:

$$egin{aligned} sk_A &= (m_A, n_A) \ pk_A &= (E_A, \phi_A(P_B), \phi_A(Q_B)) \ sk_B &= (m_B, n_B) \ pk_B &= (E_B, \phi_B(P_A), \phi_B(Q_A)) \end{aligned}$$

PK Exchange: After Alice and Bob successfully complete their key generation, they perform the following over an insecure channel:

- Alice sends Bob $(E_A, \{\phi_A(P_B), \phi_A(Q_B)\})$
- Bob sends Alice $(E_B, \{\phi_B(P_A), \phi_B(Q_A)\})$

Again, we remind the reader that we will show how curves such as E_A and E_B can be represented efficient and compactly in a computing environment when we come to our section on implementations of isogeny based systems (2.6).

SecAgr(sk_1,pk_2): After reception of Bob's tuple, Alice computes the isogeny $\phi'_A: E_B \to E_{AB}$ and Bob acts analogously. Alice and Bob then arrive at the equivalent image curve:

$$E_{AB} = \phi_A'(\phi_B(E_0)) = \phi_B'(\phi_A(E_0)) = E_0/\langle [m_A]P_A + [n_A]Q_A, [m_B]P_B + [n_B]Q_B \rangle$$

From this they can derive their shared secret k as the common j-invariant of E_{AB} .

We have included a graphical illustration of the entire SIDH key exchange process in Figure 2.5, wherein solid lines denote private computations, and dashed lines denote information sent over an insecure channel.



Figure 2.5: SIDH key exchange between Alice & Bob [FJP12]

2.3.2 Zero-Knowledge Proof of Identity

Recall the earlier discussed notion of an identification scheme. A canonical identification scheme $\Pi_{SID} = (\mathbf{KeyGen}, \mathbf{Prove}, \mathbf{Verify})$ can be derived somewhat analogously to the SIDH protocol, and is outlined in the original work of De Feo et al.

Say Bob has derived for himself the key pair (sk_B, pk_B) with $sk_B = \{m_B, n_B\}$ and $pk_B = E_B = E_0/\langle [m_B]P_B + [n_B]Q_B\rangle$ in relation to the public parameters E_0 and $\ell_B^{e_B}$. With E_0 and E_B publicly known, Π_{ZKPoI} revolves around Bob trying to prove to Alice that he knows the generator for E_B without revealing it.

To achieve this, Bob internally mimicks an execution of the key exchange protocol Π_{SIDH} with an arbitrary "random" entity Randall.

KeyGen: Key generation is performed exactly as in Π_{SIDH} , the only difference being that in Π_{ZKPoI} only the prover (Bob, in our example,) needs to generate a keypair.

Commitment: Bob generates a random point $R \in E_0[\ell_A^{e_A}]$ $(R = [m_R]P_A + [n_R]Q_A)$ along with the corresponding isogenies necessary to compute the diagram below in full (if Alice were acting as the prover in Π_{ZKPoI} , then she would choose $R \in E_0[\ell_B^{e_B}]$). Bob sends his commitment com as $(com_1, com_2) = (E/\langle R \rangle, E/\langle B, R \rangle)$ to Alice.



Response: Alice chooses a bit b at random and sends her challenge ch = b to Bob.

Prove(sk, ch): If Alice's challenge bit ch = 0 then Bob reveals the isogenies ψ_R and ψ'_R (to do this, he can simply reveal the generators of the kernels of ψ_R and ψ'_R ; R and $\phi_B(R)$ respectively). This proves he knows the information necessary to form a shared secret

with Randall if and only if he happens to know the private key $B = \{[m_B]P_B + [n_B]Q_B\}$. If ch = 1, Bob reveals the isogeny ϕ'_B . This proves that Bob knows the information necessary to form a shared secret with Randall if and only if he knows Randall's secret key R.

In the following two graphs, bold arrows are used to indicate the information revealed by Bob. The graph on the left corresponds to Bob's actions when ch = 0, the graph on the right shows the information revealed when ch = 1.

$$E_{0} - \cdots - E_{0}/\langle B \rangle \qquad E_{0} - \cdots - E_{0}/\langle B \rangle$$

$$\psi_{R} \qquad \psi_{R} \qquad$$

Note that Bob cannot at once reveal all of the information necessary to convince Alice that he knows B. If he reveals R, $\phi_B(R)$, and ϕ'_B all in one go, he incidentally reveals his secret key $B = [m_B]P_B + [n_B]Q_B$. This is because Bob reveals ϕ'_B by revealing the generator of $ker(\phi'_B)$, namely:

$$(B,R) = ([m_B]P_B + [n_B]Q_B, [m_R]P_A + [n_R]Q_A)$$

How Π_{ZKPoK} handles this is by having Bob and Alice run **Prove()** and **Verify()** for λ iterations, with a different (com, ch, resp) transcript generated for every instance. This way, if Bob is able to provide a resp that satisfies Alice's ch for every iteration, she can be sufficiently confident that Bob has knowledge of B.

Verify(pk, com, ch): Like the proving procedure, verification is a conditional function depending on the value of b:

- if ch = 0: return 1 if and only if R and $\phi_B(R)$ have order $\ell_A^{e_A}$ and generate the kernels of isogenies from $E_0 \to E_0/\langle R \rangle$ and $E_0/\langle B \rangle \to E_0/\langle B, R \rangle$ respectively.
- if ch = 1: return 1 if and only if $\psi_R(B)$ has order $\ell_B^{e_B}$ and generates the kernel of an isogeny over $E_0/\langle R \rangle \to E_0/\langle B, R \rangle$.

This scheme constitutes what is known in the literature as a zero knowledge proof of identity. It is referred to as such because Alice, acting as the verifier, does not gain any information about Bob's secret key sk.

2.4 Fiat-Shamir Construction

The Fiat-Shamir construction (also frequently referred to as the Fiat-Shamir transform, or Fiat-Shamir hueristic,) is a high-level technique for transforming a canonical identification scheme into a secure signature scheme.

The construction is rather simple. The idea is to first transform a given interactive identification protocol Π_{ID} into a non-interactive identification protocol. To achieve this, instead of allowing input from the verifier \mathcal{V} , we have our prover \mathcal{P} generate the challenge

ch by itself. In order for the verifier to be able to check that ch was generated honestly, we define ch = H(com), where H is some secure hash function. If we model H as a random oracle, H(com) is assumed truly random; from this it can be shown that it is just as difficult for an impersonator of \mathcal{P} to find an accepting transcript (com, H(com), resp) as it would be for them to successfully impersonate \mathcal{P} in Π_{ID} .

Now that we've paired Π_{ID} with H to achieve a non-interactive identification scheme Π_{NID} , we need only to factor in some message m from \mathcal{P} to have constructed a signature scheme Π'_{ID} . This can be achieved by including m in our calculation of the challenge: ch = H(com, m). Therefore, given theorem 1, if (com, H(com), resp) is an accepting transcript of Π_{NID} , then (com, H(com, m), resp) is a secure signature for the message m. Of course, because H(com, m) can be constructed by any passively observing party, it is redundant to include; and so (com, resp) constitutes a valid signature for m. A proof of theorem 1 can be found in [Kat10]. The security of the Fiat-Shamir construction was first proven by Pointcheval & Stern in [PS96].

Theorem 1 (Fiat-Shamir Security). Let Π_{ID} = (KeyGen, Commit, Prove, Verify) be a canonical identification scheme that is secure against a passive attack. Then, if H is modeled as a random oracle, the signature scheme Π'_{ID} that results from applying the Fiat-Shamir transform to Π_{ID} is classically existentially unforgeable under an adaptive chosen-message attack.

We will write $\mathbf{FS}(\Pi)$ to denote the result of applying the Fiat-Shamir transformation to some identification protocol Π .

2.4.1 Unruh's Post-Quantum Adaptation

In 2014, Ambainis et al. showed in [ARU14] that classical security proofs for "proof of knowledge" protocols are insecure in the quantum setting. This is due to a technique used in the proof of FST's security whereby the random oracle is subject to "rewinding": the proof simulates multiple runs of FST with different responses from the random oracle [ARU14].

Following this insight, Unruh proposed in [Unr14] a construction based off that of Fiat & Shamir which he proved to be secure in both the classical and quantum random oracle models.

Unruh's construction demands a small addition to the proof and verification procedures. In **Prove**, for every possible challenge value $ch_0, ch_1, ..., ch_n$, Unruh's construction demands that a hash of the corresponding responses $resp_0, resp_1, ..., resp_n$, along with the possible challenge values themselves, be included as input to the hash function H computing the actual challenge. While Unruh originally presented this technique in a generalized setting with n possible challenge values, In figure ?? we detail a version that assumes there are only two possible challenge values. This is done in an attempt to more closely reflect the zero-knowledge proof of identity scheme presented in [FJP12].

The construction is given in the form of two procedures: $\mathbf{Prove_{Un}}$ and $\mathbf{Verify_{Un}}$. Given the proving procedure \mathbf{P}_{Π} of some canonical identification scheme Π , $\mathbf{Prove_{Un}}$ can be constructed and forms the basis for the \mathbf{Sign} procedure of a quantum-safe signature scheme $\mathbf{Un}(\Pi)$. Analogously, given the verification procedure $\mathbf{V}_{\Pi} \in \Pi$, $\mathbf{Verify_{Un}}$ details the outline of signature verification in $\mathbf{Un}(\Pi)$.

Similar to above, we will write $\mathbf{Un}(\Pi)$ to denote the result of applying Unruh's construction to some identification protocol Π .

```
Algorithm 1 – Prove<sub>Un</sub>(P<sub>II</sub>)

1: if User = Alice then

2: Pick a random point S of order \ell_A^{e_A}

3: if User = Bob then

4: Pick a random point S of order \ell_B^{e_B}

5: Compute the isogeny \phi : E \to E/\langle S \rangle

6: pk \leftarrow (E/\langle S \rangle, \phi(P_{User}), \phi(Q_{User}))

7: sk \leftarrow S

8: return (sk, pk)
```

```
Algorithm 2 - Verify<sub>Un</sub>(V<sub>II</sub>)

1: if User = Alice then

2: Pick a random point S of order \ell_A^{e_A}

3: if User = Bob then

4: Pick a random point S of order \ell_B^{e_B}

5: Compute the isogeny \phi : E \to E/\langle S \rangle

6: pk \leftarrow (E/\langle S \rangle, \phi(P_{User}), \phi(Q_{User}))

7: sk \leftarrow S

8: return (sk, pk)
```

2.5 Isogeny Based Signatures

Since publication of the SIDH suite, there have been several attempts at providing authentication schemes using the same primitives. The post-quantum community had demonstrated undeniable signatures [JS14], designated verifier signatures [STW12], and undeniable blind signatures [SC16] all within the framework of isogeny based systems. It was not until the work of Yoo et al. (([YAJ+12])), however, that an isogeny based protocol for general authentication was shown as demonstrably secure. This protocol, particularly its C implementation, is where we have decided to focus our efforts.

Now that we've seen the zero-knowledge proof of identity (ZKPoI) from [FJP12] as well as Unruh's quantum-safe Fiat-Shamir adaption, we have presented all of the material necessary for an indepth analysis of the isogeny based signature scheme presented by Yoo et al. The signature protocol, which we'll denote as Σ' , is an application of Unruh's construction to the SIDH ZKPoI. In this section we will refer to the SIDH ZKPoI as Σ .

 Σ' is defined in the traditional manner, by a tuple of algorithms for key generation, signing, and verifying: $\Sigma' = (\mathbf{KeyGen}(), \mathbf{Sign}(), \mathbf{Verify}())$. $\mathbf{KeyGen}()$ in Σ' is defined identically to the key generation found in SIDH key exchange. $\mathbf{Sign}()$ and $\mathbf{Verify}()$ are defined by applying Unruh's transformation to $\mathbf{Prove}()$ and $\mathbf{Verify}()$.

For our discussion of the signature scheme, we will make use of the naming conventions used in Section 2.3. That is, we will discuss Σ' as occurring between entities Bob and Alice, with Bob imitating the role of an arbitrary third party Randall during **Sign**.

The public parameters used in Σ' are the same as outlined above for all of the protocols found in [FJP12]. Namely, we have $p = \ell_A^{e_A} \ell_B^{e_B} \cdot f \pm 1$ where $\ell_A^{e_A} = 2$, $\ell_B^{e_B} = 3$, and f is a cofacter such that p is prime. We also set as parameter the curve E such that $\#(E(F_{p^2})) = (\ell_A^{e_A} \ell_B^{e_B})^2$. And again, we include the sets of points (P_A, Q_A) and (P_B, Q_B) generating $E[\ell_A^{e_A}]$ and $E[\ell_B^{e_B}]$ respectively. We have chosen E over the previously used E_0 simply for ease of notation.

2.5.1 Algorithmic Definitions

It will be useful for us to outline in more detail the procedures of Σ' , at the very least to ease the transition into our discussion of the C implementation. In this subsection we will look at isogeny-level algorithmic definitions for **KeyGen()**, **Prove()**, and **Verify()**, and then look at how these procedures can be expressed in terms of the procedures of Π_{SIDH} .

KeyGen(λ , User): As previously mentioned, key generation in Σ' is identical to Σ :**KeyGen**(λ), which in turn is identical to Π_{SIDH} :**KeyGen**(λ). We've included a parameter User equaling either Alice or Bob – this denotes whether the user running the procedure uses blue or red constants. We've also obfuscated the lower level details in regards to how points are generated and how isogenies can be constructed. We write P_{User} and Q_{User} for P_A & Q_A or P_B & Q_B , depending on User. The result is the following:

Algorithm 3 – KeyGen(λ , User)

```
1: if User = Alice then
2: Pick a random point S of order \ell_A^{e_A}
3: if User = Bob then
4: Pick a random point S of order \ell_B^{e_B}
5: Compute the isogeny \phi : E \to E/\langle S \rangle
6: pk \leftarrow (E/\langle S \rangle, \phi(P_{User}), \phi(Q_{User}))
7: sk \leftarrow S
8: return (sk, pk)
```

Transcribing this to the procedures of Π_{SIDH} we arrive (quite trivially) at:

```
Algorithm 4 – KeyGen(\lambda, User)

1: (sk, pk) \leftarrow \Pi_{SIDH}:KeyGen(\lambda, User)
2: return (sk, pk)
```

For $\operatorname{Sign}(sk, m)$ and $\operatorname{Verify}(pk, m, \sigma)$ we assume Bob to be the signer and Alice to be the verifier. Consequently, we will write the signer's key pair (sk, pk) as (B, ϕ_B) . Algorithms for which the roles are reversed can be constructed simply by replacing red constants with their blue correspondents, and vice-versa.

 $\mathbf{Sign}(sk, m)$: The sign procedure, as a consequence of the Unruh construction, makes use of two random oracle functions \mathbf{H} and \mathbf{G} . In the sign algorithm below, make note of how Bob computes both commitments and their corresponding responses for every

iteration i before he computes the challenge values (the bits of J). He then uses the 2λ bits of J to decide which responses to include in σ .

```
Algorithm 5 – Sign(sk = B, m)
 1: for i = 1..2\lambda do
           Pick a random point R of order \ell_A^{e_A}
 2:
           Compute the isogeny \psi_R: E \to E/\langle R \rangle
 3:
           Compute the isogeny \phi_B': E/\langle B \rangle \to E/\langle B, R \rangle
 4:
           (E_1, E_2) \leftarrow (E/\langle R \rangle, E/\langle R, B \rangle)
 5:
           com_i \leftarrow (E_1, E_2)
 6:
           ch_{i,0} \leftarrow_R \{0,1\}
 7:
           ch_{i,1} \leftarrow 1 - ch_{i,0}
 8:
           (resp_{i,0}, resp_{i,1}) \leftarrow ((R, \phi_B(R)), \psi_R(B))
 9:
           if ch_{i,0} = 1 then
10:
                 Swap(resp_{i,0}, resp_{i,1})
11:
           h_{i,i} \leftarrow \mathbf{G}(resp_{i,i})
12:
13: J_1 \parallel ... \parallel J_{2\lambda} \leftarrow \mathbf{H}(\phi_B, m, (com_i)_i, (ch_{i,j})_{i,j}, (h_{i,j})_{i,j})
14: return \sigma \leftarrow ((com_i)_i, (ch_{i,j})_{i,j}, (h_{i,j})_{i,j}, (resp_{i,J_i})_i)
```

If we write out **Sign()** using the Π_{SIDH} API, we see that the only real computation is being performed by **KeyGen()** and **SecAgr()**, and our two random oracles **H()** and **G()**. The rest of the algorithm is merely organizing the information we've generated into the transcript (com, ch, resp) and then finally into σ .

```
Algorithm 6 – Sign(sk = B, m) via \Pi_{SIDH}
  1: for i = 1..2\lambda do
            (R, \psi_R) \leftarrow \Pi_{\text{SIDH}}: \mathbf{KeyGen}(\lambda, Alice)
  2:
             \phi_B': E/\langle B \rangle \to E/\langle B, R \rangle \leftarrow \Pi_{\text{SIDH}}: \mathbf{SecAgr}(B, \psi_R)
 3:
            (E_1, E_2) \leftarrow (E/\langle R \rangle, E/\langle B, R \rangle)
  4:
            com_i \leftarrow (E_1, E_2)
  5:
 6:
            ch_{i,0} \leftarrow_R \{0,1\}
            ch_{i,1} \leftarrow 1 - ch_{i,0}
  7:
            (resp_{i,0}, resp_{i,1}) \leftarrow ((R, \phi_B(R)), \psi_R(B))
 8:
            if ch_{i,0} = 1 then
 9:
                  Swap(resp_{i.0}, resp_{i.1})
10:
            h_{i,j} \leftarrow \mathbf{G}(resp_{i,j})
11:
12: J_1 \parallel ... \parallel J_{2\lambda} \leftarrow \mathbf{H}(\phi_B, m, (com_i)_i, (ch_{i,j})_{i,j}, (h_{i,j})_{i,j})
13: return \sigma \leftarrow ((com_i)_i, (ch_{i,j})_{i,j}, (h_{i,j})_{i,j}, (resp_{i,J_i})_i)
```

Verify(pk, m, σ): Alice begins her execution of Verify() where Bob ended his execution of Sign(), with the computation of J. Alice then knows at each iteration what check to perform on Bob's response, based on a conditional branch. You will notice that Bob's secret key B occurs in the negative path of this branch; this is not a security concern because it is actually the point $\psi_R(B)$ that is communicated in σ , from which B cannot be recovered.

Algorithm 7 - Verify $(pk = \phi_B, m, \sigma)$

```
1: Parse ((com_i)_i, (ch_{i,j})_{i,j}, (h_{i,j})_{i,j}, (resp_i)_i) \leftarrow \sigma
2: J_1 \parallel ... \parallel J_{2\lambda} \leftarrow \mathbf{H}(\phi_B, m, (com_i)_i, (ch_{i,j})_{i,j}, (h_{i,j})_{i,j})
 3: for i = 0..2\lambda do
         check h_{i,J_i} = G(resp_i)
 4:
         if ch_{i,J_i} = 0 then
 5:
 6:
              Parse (R, \phi_B(R)) \leftarrow resp_i
              check (R, \phi_B(R)) have order \ell_A^{e_A}
 7:
              check R generates the kernel of the isogeny E \to E_1
 8:
 9:
              check \phi_B(R) generates the kernel of the isogeny E/\langle B \rangle \to E_2
         else
10:
              Parse \psi_R(B) \leftarrow resp_i
11:
              check \psi_R(B) has order \ell_B^{e_B}
12:
13:
              check \psi_R(B) generates the kernel of the isogeny E_1 \to E_2
14: if all checks succeed then
         return 1
15:
16: else
17:
         return 0
```

What we are checking for in the verification process is whether or not Bob and Randall performed an honest and valid key exchange. And so, if the challenge bit is 0, we can use SIDH key generation to determine that R and ψ_R are a valid key pair and then run SIDH secret agreement with R and Bob's public key ϕ_B to confirm that it properly executes outputting an isogeny with kernel generated by $\phi_B(R)$. If the challenge bit is 1, we can run an instance of SIDH secret agreement to verify that $\psi_R(B)$ generates the kernel of an isogeny with domain E_1 and co-domain E_2 (refer again to the diagrams outlining **Prove** of section 2.3.2).

These observations are formalized in Algorithm 6, where we rewrite Σ' :Verify() in terms of Π_{SIDH} procedure calls. Note, in line 10 of Algorithm 6, the call Π_{SIDH} :SecAgr($\psi_R(B)$, ψ_R). It should be noted that $\psi_R(B)$ is not the proper secret key input used by Bob in Sign(), but we will see in the section to follow how we can use $\psi_R(B)$ in the C implementation of SecAgr to perform our verification (without compromising Bob's secret key B).

```
Algorithm 8 – Verify(pk = \phi_B, m, \sigma) via \Pi_{\text{SIDH}}
 1: Parse ((com_i)_i, (ch_{i,j})_{i,j}, (h_{i,j})_{i,j}, (resp_i)_i) \leftarrow \sigma
 2: J_1 \parallel ... \parallel J_{2\lambda} \leftarrow \mathbf{H}(\phi_B, m, (com_i)_i, (ch_{i,j})_{i,j}, (h_{i,j})_{i,j})
 3: for i = 0..2\lambda do
         check h_{i,J_i} = G(resp_{i,J_i})
 4:
         if ch_{i,J_i} = 0 then
 5:
 6:
              Parse (R, \phi_B(R)) \leftarrow resp_{i,J_i}
              check (R, \psi_R) is a valid output of \Pi_{SIDH}: KeyGen(\lambda, Alice)
 7:
              check that \Pi_{\text{SIDH}}:SecAgr(R, \phi_B) successfully outputs an isogeny with co-
 8:
     domain E_2
         else
 9:
10:
              Parse \psi_R(B) \leftarrow resp_{i,J_i}
              check that \Pi_{\text{SIDH}}:SecAgr(\psi_R(B), \psi_R) successfully outputs an isogeny with
11:
     co-domain E_2
12: if all checks succeed then
         return 1
13:
14: else
```

2.6 Implementations of Isogeny Based Cryptographic Protocols

Having now introduced all of the background material necessary for understanding SIDH and the isogeny based signature scheme in detail, we will investigate the portions of the SIDH C library which are relevent to our contributions.

The SIDH C library, written by a research team at Microsoft Research, was released in 2016 alongside an article titled Efficient Algorithms for Supersingular Isogeny Diffie-Hellman (see [CLN16]). The article in question details several adjustments to the algorithms and data-representations outlined in [FJP12], leading to improved performance and key-sizes. Their library (which we will henceforth refer to as SIDH_C) consists of C and assembly implementations of the algorithms outlined in [CLN16]. Much of these functions are tailored to a specific set of parameters allowing for increased performance. The library presents 128-bit quantum security and 192-bit classical security key exchange up to 2.9 times faster than any previous isogeny-based key-exchange system. We will look at some of the details of SIDH_C below.

Before proceeding, it may be advisable to briefly review the section on notation (??) if one has not already.

2.6.1 Parameters & Data Representation

15:

return 0

Parameters. SIDH_C operates over the underlying basefield \mathbb{F}_p where $p = \ell_A^{e_A} \cdot \ell_B^{e_B} - 1$, with $\ell_A = 2$, $\ell_B = 3$, $\ell_A = 372$, and $\ell_B = 239$, giving p a bitlength of 751. Now, recall the Montgomery representation of a curve:

$$By^2 = Cx^3 + Ax^2 + Cx$$

SIDH_c uses the public parameter curve E defined in Montgomery form with A=0, B=1, and C=1. The point pairs (P_A,Q_A) and (P_B,Q_B) , generating $E[\ell_A^{e_A}]$ and

 $E[\ell_B^{e_B}]$ respectively, are hard-coded as an array of bytes. These parameters (including related data such as the bitlength of certain constants) are stored in the struct type CurveIsogenyStaticData under the variable name SIDHp751. This struct, along with many other SIDH_C data types and representations, will be outlined in the coming subsection.

One priority of the parameter choices found in SIDHp751 was to approach $\ell_A^{e_A} \approx \ell_B^{e_B}$. This attempted at balancing $\ell_A^{e_A}$ and $\ell_B^{e_B}$ helps to ensure two things: first, that no side of the key exchange is any easier to attack than the other, and second, that the cost of computation is split evenly between parties. This constraint had to be compromise with the primary security concern: that p must have a bit-length providing sufficient classical and quantum security.

Data Structures. There are several custom-defined data structures that are integral to SIDH_c. Below, we will briefly cover the ones which are likely to arise in our discussion:

Field elements

- felm_t buffer of bytes representing elements of \mathbb{F}_p .
- f2elm_t pair of felm_t representing elements of \mathbb{F}_{p^2} .

Elliptic curve points

- point_affine an f2elm_t x and an f2elm_t y representing a point in affine space.
- point_proj an f2elm_t X and an f2elm_t Z representing a point as projective XZ Montgomery coordinates.
- point_full_proj f2elm_t elements X, Y, and Z representing a point in projective space.
- point_basefield_affine an felm_t x and an felm_t y representing a point in affine space over the base field.
- point_basefield_proj an felm_t X and an felm_t Z representing a point as projective XZ Montgomery coordinates over the base field.

Cryptographic structures

publickey_t - three f2elm_ts representing a public key.
 publickey_t[0] = user's private isogeny applied to the other party's generator P_x
 publickey_t[1] = user's private isogeny applied to the other party's generator Q_x
 publickey_t[2] = user's private isogeny applied to P_x - Q_x

Curve structures

- CurveIsogenyStruct Structure containing all necessary public parameter data.
- CurveIsogenyStaticData The same as CurveIsogenyStruct, but with buffer sizes fixed for SIDHp751.

The reader may note that publickey_t does not contain any information defining the user's co-domain curve $E/\langle S \rangle$ (with S as the users secret key). It just so happens that in Π_{SIDH} key exchange, the curves $E/\langle A \rangle$ and $E/\langle B \rangle$ are simply intermediary steps (useful for conceptualizing the protocol) and not necessary for computing the shared secret $j(E_{AB})$.

Also worth noting is the lack of a specific data structure for representing curves. As it turns out, curves within Π_{SIDH} can be distinctly represented by their A value alone. As we are working with curves defined over \mathbb{F}_{p^2} , we have $A \in \mathbb{F}_{p^2}$ and thus we can succinctly represent any curve with a single f2elm_t.

2.6.2 SIDH_C Design Decisions

The following are, at a high-level, the algorithmic improvements upon Π_{SIDH} as outlined in [CLN16]. Costella et al. do make additional contributions in their paper, however we will discuss only those contributions which pertain to the performance of SIDH.

Projective Space Arithmetic. As is common in ECC, a vast majority of the procedures of SIDH_C operate over elliptic curve points which are defined over projective space (recall Section 2.2.2). This widely-deployed technique is used to avoid the substantial cost of field element inversions (computing x^{-1} for some element $x \in \mathbb{F}_{p^2}$). This means the majority of our calculations are performed over point_proj structures using Montgomery arithmetic (Section 2.2.1) and converted to point_affine when necessary. This general design strategy is highly related to our first contribution, which will be elaborated upon in Section 3.

In addition to traditional point-wise projective arithmetic, Costella et al. showed that isogeny arithmetic can also be carried out in this space. By performing isogeny arithmetic in the projective space, the number of \mathbb{F}_{p^2} inversions in Π_{SIDH} :**KeyGen** and Π_{SIDH} :**SecAgr** can be reduced to 1 and 2, respectively.

Key Representation. Recall the origin of an Π_{SIDH} private key (m, n): the goal is to randomly select a generator of the torsion group $E[\ell_A^{e_A}]$ (or $E[\ell_B^{e_B}]$ for Bob). It is noted in [FJP12] that any generator of the required torsion group is sufficient. It is also noted that m, unless equal to the order of the torsion group, is invertible. Because of this, Alice, for example, can simple compute $R = P_A + [m^{-1}n]Q_A$, thus enabling secret keys to be stored as a single \mathbb{F}_{p^2} element (which is referred to as m). This technicality has been implemented in SIDH_C, which both saves on storage as well as offers a means for generating secret keys that is more efficient than the trivial scalar multiplication and point-wise addition approach to computing [m]P + [n]Q.

Tailor-made Montgomery Multiplication. The parameters of a default SIDH_C execution, stored in SIDHp752, support efficient arithmetic and grant access to a variety of modular arithmetic optimizations. These optimizations inlude esfs,esdfsef,sefsef.. Moreover, Costella et al. supply a modified version of the Montgomery multiplication algorithm which, when performing over the class of curves outlined by their set of parameters, yields faster modular arithmetic.

2.6.3 Key Exchange & Critical Functions

There are 3 central modules (C files) in $SIDH_C$, all dealing with different levels of abstraction in the Π_{SIDH} protocol. Figure 2.6 illustrates the relationship between these modules and the abstraction levels of isogeny based key exchange.

Operating at the lowest abstraction level is the module fpx.c, wherein functions for manipulating \mathbb{F}_p and \mathbb{F}_{p^2} elements are defined. One level up from fpx.c we have $ec_isogeny.c$, containing functions pertaining to elliptic curves and point arithmetic (such as $j_inv(...)$ for computing the j-invariant of a curve and $secret_pt(...)$ for computing a users secret point S given their private key m). The final, highest abstraction-level module we will discuss is kex.c. kex.c contains the protocol-level functions for performing Π_{SIDH} , namely $KeyGeneration_A(...)$ and $KeyGeneration_B(...)$ for generating Alice and Bob's private and public keys, as well as $SecretAgreement_A(...)$ and $SecretAgreement_B(...)$ for completing the secret agreement from both sides of the key exchange.



Figure 2.6: Relationship between Π_{SIDH} & SIDH_c modules

For functions defined in fpx.c the notational practice is to prepend function names with either fp or fp2, signifying whether the function is defined for elements of \mathbb{F}_p or \mathbb{F}_{p^2} . Additionally, it is common to append mont to the name of functions which utilize Montgomery arithmetic, and thus expect elements in Montgomery representation. Functions in fpx.c are largely defined by byte and memory arithmetic, with the exception of slightly higher-level functions (such as field element inversion, fpinv751_mont(...)) which are defined in terms of other fpx.c functions. Furthermore, for efficiency, functions of fpx.c are defined as __inline when applicable.

In addition to the fpx.c functions we've outlined in Figure 2.6.3 there are of course definitions for addition, copying elements, retrieving the zero element, Montgomery multiplication, squaring, and so on and so forth.

ec_isogeny.c functions are defined almost exclusively in terms of fpx.c functions, with a few occurances of internal function calling. Functions in this module that are significant to our our work are briefly summerized in Figure 2.6.3. The implementation specifics of most other ec_isogeny.c functions are not critical to our work, and so have been excluded. The design and efficiency of these algorithms do, however, have a rich background and can be further read about in [FJP12] and [CLN16].

The key exchange procedures found in kex.c are composed entirely of calls to fpx.c and ec_isogeny.c functions, modulo some basic branching logic. All of the functions

fpx.c functions

Function	Input	Output
to_fp2mont	f2elm_t a	f2elm_t ma
Converts an \mathbb{F}_{p^2} element		
to Montgomery representation		
from_fp2mont	f2elm_t ma	f2elm_t a
Converts an \mathbb{F}_{p^2} element		
from Montgomery representation		
to regular form		
fp2inv751_mont_bingcd	f2elm_t a	$f2elm_t a^{-1}$
performs non-constant		
time inversion of		
a \mathbb{F}_{p^2} element		
fp2inv751_mont	f2elm_t a	$f2elm_t a^{-1}$
performs constant		
time inversion of		
a \mathbb{F}_{p^2} element		

ec_isogeny.c functions

Function	Input	Output
j_inv	f2elm_t A	f2elm_t jinv
computes the j-invariant	f2elm_t C	
of a curve with represented		
in Montgomery form with A and C		
secret_pt	point_basefield P	point_proj R
generates the secret	$ ext{digit}_{ ext{-}} ext{t}$ m	
point R from	SIDHp751	
secret key m	int AliceOrBob	
inv_3_way	f2elm_t z1	$f2elm_t z1^{-1}$
performs simultaneous inversion	f2elm_t z2	$ exttt{f2elm_t} exttt{z2}^{-1}$
of three elements	f2elm_t z3	$f2elm_t z3^{-1}$
${ t inv}_4$ _way	f2elm_t z1	$f2elm_t z1^{-1}$
performs simultaneous inversion	f2elm_t z2	$ exttt{f2elm_t} exttt{z2}^{-1}$
of 4 elements	f2elm_t z3	$ exttt{f2elm_t} exttt{z3}^{-1}$
	f2elm_t z4	$ exttt{f2elm_t} exttt{z4}^{-1}$
generate_2_torsion_basis	f2elm_t A	point_full_proj R1
constructs a basis $(\{R1, R2\})$	SIDHp751	point_full_proj R2
generating $E[\ell_A^{e_A}]$		
generate_3_torsion_basis	f2elm_t A	point_full_proj R1
constructs a basis $(\{R1, R2\})$	SIDHp751	point_full_proj R2
generating $E[\ell_B^{e_B}]$		

kex.c functions

Function	Input	Output	
KeyGeneration_A	unsigned char* privateKeyA	unsigned char* privateKeyA	
performs key generation	bool generateRandom	unsigned char* publicKeyA	
for Alice			
KeyGeneration_B		unsigned char* privateKeyB	
performs key generation		unsigned char* publicKeyB	
for Bob			
SecretAgreement_A	unsigned char* privateKeyA	unsigned char* sharedSecretA	
computes the shared secret	unsigned char* publicKeyB	point_proj kerngen	
from Alice's perspective	point_proj kerngen		
SecretAgreement_B	unsigned char* privateKeyB	unsigned char* sharedSecretB	
computes the shared secret	unsigned char* publicKeyA	point_proj kerngen	
from Bob's perspective	point_proj kerngen		

from this module are relevant to our work - we provide quick debriefings of these functions in Figure 2.6.3.

The reader may note that, in Figure 2.6.3, privateKeyA (in KeyGeneration_A) and kerngen (in both secret agreements) appear as both inputs and outputs. This is not a mistake. In KeyGeneration_A, if generateRandom = false is passed as an input, then privateKeyA is expected to be set, and the corresponding public key is computed. In secret agreement, if kerngen is set to null then the algorithm proceeds normally. If it is set to a valid point, however, it can be used in place of a secret key input (which in such a case is expected to be null). Both of these details are critical to the design of signature functions as they are described below.

2.6.4 Signature Layer

Yoo et al. provided, along with their publication of [YAJ $^+$ 12], an implementation of their signature scheme as a fork to SIDH $_c$. All of their functions are written specifically for an instance of Σ' where the signer is assuming the B role (meaning that Randall assumes the A role), but their algorithms could be trivially modified to provide versions supporting a signer in the A role. Their contributions to the SIDH $_c$ codebase come in the form of the functions listed below.

Function	Input	Output
isogeny_keygen generates the signers key pair		unsigned char* privateKeyB unsigned char* publicKeyB
isogeny_sign produces a signature for a message sign_thread performs a single iteration of the for-loop in Sign	privateKey publicKey message m	Signature sig
isogeny_verify checks the validity of a signature verify_thread	Signature sig	true or false
performs a single iteration of the for-loop in Verify		

To begin, isogeny_keygen has a trivial definition; KeyGeneration_B is called and populates the signer's public and private keys. isogeny_keygen returns the success status of the call to KeyGeneration_B.

In their original fork of SIDH_c, Yoo et al. included these functions in the file kex_tests.c. This file was originally intended for testing the functions of kex.c, and so our fork of the library has placed the signature functions in a new file SIDH_signature.c. We have also included a file sig_tests.c for testing the contents and performance of SIDH_signature.c functions.

If we transcribe the procedures Σ' :**Sign** and Σ' :**Verify** (as described in Section ??) to the language of the SIDH_C API, we have in essence the procedures Sign and Verify given by Algorithms 9 and 10 respectively.

```
Algorithm 9 – Sign(sk_B, m)
```

```
1: for i = 1..2\lambda do

2: (sk_R = R, pk_R) \leftarrow \text{KeyGeneration\_A(NULL, true)}

3: (E/\langle B, R \rangle, \psi_R(B)) \leftarrow \text{SecretAgreement\_B}(sk_B, pk_R, \text{NULL})

4: (E_1, E_2) \leftarrow (E/\langle R \rangle, E/\langle B, R \rangle)

5: (\text{com}[i]_0, \text{com}[i]_1) \leftarrow (E_1, E_2)

6: (\text{resp}[i]_0, \text{resp}[i]_1) \leftarrow (R, \psi_R(B))

7: h[i] \leftarrow \text{keccak}(\text{resp}[i]_0) | \text{keccak}(\text{resp}[i]_1)

8: J_1 \parallel ... \parallel J_{2\lambda} \leftarrow \text{keccak}(\text{com}, m, h)

9: \text{return } \sigma \leftarrow ((\text{com}_i)_i, (\text{ch}_{i,j})_{i,j}, (h_i)_i, ((\text{resp})[J_i])
```

I'm shook.

Algorithm 10 – Verify($pk = \phi_B$, m, σ) 1: $J_1 \parallel ... \parallel J_{2\lambda} \leftarrow \texttt{keccak(com, } m, h)$ 2: **for** i = $0..2\lambda$ **do** $\mathbf{check}\ h[i] = \mathtt{keccak}(\mathtt{resp}[i]_0) \,|\, \mathtt{keccak}(\mathtt{resp}[i]_1)$ 3: 4: if $J_i = 0$ then 5: $R \leftarrow \text{resp}[i]_0$ $pk_R \leftarrow \text{KeyGeneration_A(R, false)}$ 6: $\mathbf{check}\ pk_{R} = \mathbf{com}[i]_{0}$ 7: $E_{RB} \leftarrow \texttt{SecretAgreement_A}(R, \phi_B, \texttt{NULL})$ 8: $\mathbf{check}\ E_{RB} = \mathbf{com}[i]_1$ 9: 10: else $\psi_R(B) \leftarrow \text{resp}[i]_1$ 11: $pk_{R} \leftarrow \text{com}[i]_{0}$ 12: $E_{BR} \leftarrow \texttt{SecretAgreement_B(NULL, } pk_R, \ \psi_R(B))$ 13: $\mathbf{check}\ E_{BR} = \mathbf{com}[i]_1$ 14: 15: if all checks succeed then 16: return 1 17: **else** 18: return 0

Outside of simply replacing Π'_{SIDH} procedure calls with SIDH_C functions, the reader may notice additional differences between Sign and Verify and their Σ' counterparts. Namely, Yoo et al. have chosen to exclude the challenge bit ch in the SIDH_C implementations of these functions, consequently excluding the conditional and Swap statement of lines 8 and 9 in Algorithm 4.

Chapter 3

Batching Operations for Isogenies

Our first contribution to the $SIDH_{C}$ codebase is the implementation and integration of a procedure for batching together many \mathbb{F}_{p^2} element inversions. This contribution is discussed in detail in the following chapter. The chapter is split into three sections: a high-level discussion of the procedure itself, the low-level details of its integration into $SIDH_{C}$, and finally, the resulting affects of this procedure on the performance of $SIDH_{C}$.

In the first section of this chapter we will detail the specifics of the partial batched inversion procedure. We will show how the procedure can be constructed by combining two techniques: a well known method for reducing a \mathbb{F}_{p^2} inversion to several \mathbb{F}_p operations, and an inversion batching technique outlined in [SB01].

As we then venture into the lower-level implementation details, we will explore how the procedure can be leveraged efficiently and intelligently in the codebase. We will take a closer look at several of the aforementioned SIDH_C functions as we illustrate some of the performance bottlenecks in the system. At this time, we will also discuss the design decisions made while implementing the partial batched inversion procedure as well as some of the function's lower-level minutiae.

We will end this chapter by taking a detailed look at the performance gains offered by the inclusion of partial batched inversions in SIDH_c. More precisely, we will be examining the effects of the procedure on the Yoo et al. signature layer. We will contrast the measured performance of our implementation with an analytical calculation of the expected improvement, and discuss the possible origins of divergent behaviour.

3.1 Partial Batched Inversions

We will now outline the procedure that is central to our first contribution. The "partial batched inversion" procedure reduces arbitrarily many $unrelated^1 \mathbb{F}_{p^2}$ inversions to a sequence of \mathbb{F}_p operations. The fact that the elements being inverted need not hold any relation will be significant to the applicability of this procedure. For brevity's sake, we will henceforth refer to this procedure as pb_i inv in the SIDH_C context, and Partial-BatchedInversion in the more general mathematical context.

As mentioned above, pb_{inv} is constructed by combining two distinct techniques. Both of these techniques improve the efficiency of computing field element inversions: the first is specific to extension fields (in our case, \mathbb{F}_{p^2} elements,) but the second is a

¹To clarify; the elements subject to these inversion must all be over the same field, but can otherwise be unrelated.

technique applicable to field element inversions in a more general setting.

We will begin with a dissection of these two techniques, starting first with the "partial" inversion technique and then looking at batched inversions. The definitions we will give for these techniques below are given at the level of field arithmetic. When we proceed to sketch pb_inv, we will offer two definitions: one in this section given at the abstraction-level of field arithmetic, and one in the proceeding section given in terms of SIDH_C syntax.

In the subsections to come, when we are working at the level of field arithmetic we will denote the first and second portions of an arbitrary $x \in \mathbb{F}_{p^2}$ as x_a and x_b respectively, where $x = x_a + x_b \cdot i$. Additionally, we may write x as $\{x_a, x_b\}$, as this more closely reflects the structure of \mathbb{F}_{p^2} elements in SIDH_C. Recall from Section 2.2.1 that both x_a and x_b are valid \mathbb{F}_p elements.

We will express the time-complexity of the coming procedures in terms of the number of underlying field operations within them. We denote the computation time for base field arithmetic with bold letters (such as **a** for \mathbb{F}_p addition), and we use bold letters accented with a "closure" bar for extension field arithmetic ($\bar{\mathbf{a}}$ for \mathbb{F}_{p^2} addition). For example, the time-complexity of some procedure P, which we might write as C_P , may look like the following:

$$C_P = 2\bar{\mathbf{a}} + x\bar{\mathbf{i}} + y\mathbf{m} + \mathbf{s}$$

Which denotes that P is a procedure composed of $2 \mathbb{F}_{p^2}$ additions, x-many \mathbb{F}_{p^2} inversions, y-many \mathbb{F}_p multiplications, and a single \mathbb{F}_p squaring. We reserve uppercase bold letters for arithmetic over elliptic curve points (such as \mathbf{A} to denote the point-wise addition operation).

3.1.1 \mathbb{F}_{p^2} Inversions done in \mathbb{F}_p

There is a simple way in which we can perform one \mathbb{F}_{p^2} inversion by means of doing several \mathbb{F}_p operations. We will begin by considering multiplicative inverses of complex numbers. Fields of the form \mathbb{F}_{q^2} for some prime q are, after all, quadratic extension fields; because of this \mathbb{F}_{p^2} arithmetic is treated, for the most part, analogously to complex number arithmetic.

Consider the complex number C = a + bi. We have that $C^{-1} = 1/(a + bi)$, from which we can rationalize the denominator like so:

$$C^{-1} = \frac{1}{(a+bi)} \cdot \frac{(a-bi)}{(a-bi)}$$
$$C^{-1} = \frac{a-bi}{(a+bi)(a-bi)}$$

Here we note that (a+bi)(a-bi) is equivalently (a^2-b^2) and so we can rewrite C^{-1} as the following:

$$C^{-1} = \frac{a - bi}{(a)^2 - (bi)^2}$$
$$C^{-1} = \frac{a - bi}{a^2 + b^2}.$$

Elements in the quadratic extension of a finite field are treated similarly, such that if we take some element $x = \{x_a, x_b\} \in \mathbb{F}_{p^2}$ for some prime p, we can equivalently represent

x as $x_a + x_b i$ and treat arithmetic on x exactly as we would for a complex number (modulo p, of course). From this we can see that x^{-1} can be defined as:

$$x^{-1} = \left\{ \frac{x_a}{x_a^2 + x_b^2}, \frac{-x_b}{x_a^2 + x_b^2} \right\}$$

Now it is clear that we can compute the multiplicative inverse of x by computing the inverse of $x_a^2 + x_b^2$ (an inversion in \mathbb{F}_p) and $-x_b$ (a relatively inexpensive operation, also in the base field). We formulate this technique in Algorithm 13, which we refer to as PartialInv.

$oxed{\mathbf{Algorithm}} \ \mathbf{11} - \mathbf{PartialInv}(x \in \overline{\mathbb{F}}_{p^2})$

- 1: $den \leftarrow x_a^2 + x_b^2$ 2: $den_{inv} \leftarrow den^{-1} \pmod{p}$
- 3: $a \leftarrow x_a \cdot den_{inv} \pmod{p}$
- 4: $b \leftarrow -(x_b) \cdot den_{inv} \pmod{p}$
- 5: $inv \leftarrow \{a, b\}$
- 6: return inv

Effectively, this procedure reduces one \mathbb{F}_{p^2} inversion to the following operations:

- 2 \mathbb{F}_p squarings line 1 of algorithm 13
- 1 \mathbb{F}_p addition line 1 of algorithm 13
- 1 \mathbb{F}_p inversion line 2 of algorithm 13
- 3 \mathbb{F}_p multiplications lines 3 & 4 of algorithm 13

Let $C_{\mathbf{PartialInv}}$ represent the performance of $\mathbf{PartialInv}$, in the format outlined above. We have

$$C_{\text{PartialInv}} = 2\mathbf{s} + \mathbf{a} + \mathbf{i} + 3\mathbf{m}$$

In some contexts, computing squares can be done more efficiently than the multiplication of two arbitrary elements. A noteworthy example of this is would be binary fields (\mathbb{F}_{2^k}) where squaring a number is equivalent to simply performing a bit-shift. However, because we are working in the quadratic extension of some prime field \mathbb{F}_p for a large prime p, we can assume that computing the square of some arbitrary element x is no more or less efficient than simply computing $x \cdot x$. With this in mind, we can further simplify A.

$$C_{\mathbf{PartialInv}} = 5\mathbf{m} + \mathbf{a} + \mathbf{i}$$

We claim that this "reduces" the computation time of field element inversion, but it may not be immediately clear that this technique is computationally favourable over any other approach to computing an \mathbb{F}_{p^2} inversion.

3.1.2 Batching Field Element Inversions

The second technique used in the composition of pb_inv reduces arbitrarily many (general) field element inversions to *one* inversion and a linearly scaling amount of multiplications in the *same* field.

This technique was outlined by Shacham and Boneh in [SB01]. Shacham and Boneh provided several techniques for improving the performance of SSL handshakes, most of which built on the earlier efforts of Amos Fiat in batching multiple RSA decryptions ([Fia89]). While somewhat related, Fiat's work admittedly is only applicable to the RSA cryptosystem, and requires additional constraints on the elements being batched.

One improvement offered by Shacham and Boneh, however, is their proposed notion of batching together divisions from across multiple unrelated SSL instances.

Suppose we want to compute the inverses of three elements $x, y, z \in F$ where F is some arbitrary field. The batched division technique allows us to reduce these three inversions to one. The technique can be organized into three phases. In the first phase, all the elements of the batch are multiplied together into one product, yielding a = xyz. We refer to this first phase as "upward-percolation". Next, we compute the inverse of a: $a^{-1} = (xyz)^{-1}$, which we refer to as the inversion phase. In the final phase, "downward-percolation", we can compute each individual element's multiplicative inverse as follows:

$$x^{-1} = a^{-1} \cdot (yz)$$

$$y^{-1} = a^{-1} \cdot (xz)$$

$$z^{-1} = a^{-1} \cdot (xy)$$

Let us analyse these phases a little more closely while we generalize to n-many elements. In the upward-percolation phase, constructing a requires n-1 multiplications; and so has a complexity of $\mathcal{O}(n)$. The inversion phase requires one field element inversion, and so has complexity of $\mathcal{O}(1)$.

If we implement the downward-percolation phase directly as outlined in the threeelement example above, computing every output requires n products each composed of n-1 multiplications. These n products are each also multiplied by a^{-1} . This multiplication by a^{-1} can be added to our n-1 inversion count resulting in n-many products composed of n multiplications; bringing the complixity of the downward-percolation phase to $\mathcal{O}(n^2)$.

We will refer to this roughly-sketched procedure as **BatchedInv**₀. Let $C_{\mathbf{BatchedInv}_0}$ denote the performance of **BatchedInv**₀ in the format outlined above. We have, then, that

$$C_{\mathbf{BatchedInv}_0} = n^2 \bar{\mathbf{m}} + (n-1)\bar{\mathbf{m}} + \bar{\mathbf{i}}.$$

This batching proceedure can be thought of as analogous to traditional time-memory tradeoff algorithms. In a general time-memory tradeoff algorithm you can continue to make some linear or polynomial (or otherwise) sacrifice of memory in order to gain some increase in performance. In the batching procedure described above we are in some sense sacrificing some marginal amount of memory to gain an increase in performance, but it is not a tradeoff that we can adjust to our liking.

There is a way, much akin to this time-memory tradeoff strategy, that we can further reduce the execution time of **BatchedInv**₀. In the upward-percolation phase, we currently store in a the product of elements $x_0 \cdot x_1 \cdot ... \cdot x_{n-1}$. Suppose instead that we store

in a an array (size n) of elements, defined in the following way:

$$a_i = \begin{cases} x_0 & i = 0\\ a_{i-1} \cdot x_i & \text{otherwise} \end{cases}$$

Equivalently, the elements of this array are

$$a_0 = x_0, \quad a_1 = x_0 \cdot x_1, \quad a_2 = x_0 \cdot x_1 \cdot x_2, \quad \dots$$

and so on and so forth up to n-1. In the inversion phase we will compute $inv = a_{n-1}^{-1}$; we are still inverting the product of all the elements, but because we have stored the value of the product at every step of the way, we can save on a significant number of operations in the downward-percolation phase.

Going into the final stage of the procedure now, we can compute x_{n-1}^{-1} simply by computing $inv \cdot a_{n-2}$. Moving forward (or backwards, technically), we peel the previously used x_{n-1}^{-1} off of inv by computing $inv := inv \cdot x_{n-1}$ and, with our updated inv, we compute $x_{n-2}^{-1} = inv \cdot a_{n-3}$. We proceed in this fashion until we reach x_0^{-1} , which (if we've been updating inv every step of the way) is simply equal to inv.

We formalize this improvement in the form of a new procedure, **BatchedInv**, which we provide a concrete definition for in Algorithm 12. In this procedure lines 1–3 implement the upward-percolation phase. Line 4 carries out the second phase: the inversion of a_{n-1} . The third and final stage, downward-percolation, occurs from lines 5 to 7.

Algorithm 12 – BatchedInv $(\{x_0, x_1, ..., x_n - 1\} \in \mathbb{F}_{p^2}^n)$

```
1: a_0 \leftarrow x_0

2: for i = 1...(n-1) do

3: a_i \leftarrow a_{i-1} \cdot x_i

4: inv \leftarrow a_{n-1}^{-1}

5: for i = (n-1)...1 do

6: x_i^{-1} \leftarrow a_{i-1} \cdot inv

7: inv \leftarrow inv \cdot x_i

8: x_0^{-1} = inv

9: return \{x_0^{-1}, x_1^{-1}, ..., x_{n-1}^{-1}\}
```

BatchedInv can be used to reduce n-many \mathbb{F}_{p^2} inversions to the following operations:

- n-1 \mathbb{F}_{p^2} multiplications line 2-3 of algorithm 12
- 1 \mathbb{F}_{p^2} inversion line 4 of algorithm 12
- 2(n-1) \mathbb{F}_{p^2} multiplications line 5-7 of algorithm 12

Let $C_{\mathbf{BatchedInv}}$ denote the performance of $\mathbf{BatchedInv}$.

$$C_{\mathbf{BatchedInv}} = 2(n-1)\bar{\mathbf{m}} + (n-1)\bar{\mathbf{m}} + \bar{\mathbf{i}}$$

= $3(n-1)\bar{\mathbf{m}} + \bar{\mathbf{i}}$

In comparing the performances of **BatchedInv** and **BatchedInv**₀, we see that $C_{\mathbf{BatchedInv}} < C_{\mathbf{BatchedInv_0}}$ holds when the following holds:

$$2(n-1)\bar{\mathbf{m}} + (n-1)\bar{\mathbf{m}} + \bar{\mathbf{i}} < n^2\bar{\mathbf{m}} + (n-1)\bar{\mathbf{m}} + \bar{\mathbf{i}}$$
$$2(n-1)\bar{\mathbf{m}} < n^2\bar{\mathbf{m}}$$
$$2(n-1) < n^2$$

And so, because n^2 is always larger than 2(n-1) for all $n \in \mathbb{R}$, **BatchedInv** outperforms **BatchedInv**₀ for every possible batch size. This can be checked, if one so wishes, by setting $n^2 = 2(n-1)$, simplifying to $n^2 - 2n + 2 = 0$, and noting that the discriminant $(2^2 - 4 \cdot 2)$ is negative.

Partial Batched Inversions 3.1.3

We have now outlined the following: **PartialInv** as a technique for computing \mathbb{F}_{p^2} inversions by means of \mathbb{F}_p arithmetic, and **BatchedInv** as a technique for batching together arbitrarily many inversion operations. We will now combine these procedures to achieve the partial batched inversion algorithm.

At first glance, an attempt to meld these two techniques together might be made in the same fashion as Algorithm 13. We denote this approach PartialBatchedInv₀.

```
Algorithm 13 – PartialBatchedInv<sub>0</sub>(\{x_0, x_1, ..., x_{n-1}\})
```

- 1: $a \leftarrow \text{upward-percolation of elements } \{x_0, x_1, ..., x_{n-1}\}$
- 2: $a^{-1} \leftarrow \mathbf{PartialInv}(a)$
- 3: $\{x_0^{-1}, x_1^{-1}, ..., x_{n-1}^{-1}\} \leftarrow \text{downward-percolation of } a^{-1}$ 4: $\mathbf{return} \ \{x_0^{-1}, x_1^{-1}, ..., x_{n-1}^{-1}\}$

If we sum the operations in $PartialBatchedInv_0$, we have the following:

- $n \mathbb{F}_{p^2}$ multiplications upward-percolation phase
- 2 \mathbb{F}_p squarings, 1 \mathbb{F}_p addition, 1 \mathbb{F}_p inversion, and 3 \mathbb{F}_p multiplications call to PartialInv(a)
- $2n \mathbb{F}_{p^2}$ multiplications downward-percolation phase

To measure the complixity in terms of field operations, denoted C_0 , we can surmize the the total operation count as:

$$C_0 = (n\bar{\mathbf{m}}) + (2\mathbf{s} + \mathbf{a} + \mathbf{i} + 3\mathbf{m}) + (2n\bar{\mathbf{m}})$$
$$C_0 = 3n\bar{\mathbf{m}} + 2\mathbf{s} + \mathbf{a} + \mathbf{i} + 3\mathbf{m}$$

Below we provide an alternative approach to building PartialBatchedInv that relies on only \mathbb{F}_p operations. Afterward, we show by simple analysis why this approach yields the better performance. This procedure is formalized in a mathematical setting in Algorithm 14. We give a precise C function definition in Section 3.2.

In Algorithm 14, a is a simple auxiliary set we use to hold the inverted \mathbb{F}_p elements. After these are all computed via the for-loop on line 8, we can reconstruct \mathbb{F}_p .

More specifically, the procedure takes us from $n \mathbb{F}_{p^2}$ inversions to:

Algorithm 14 – PartialBatchedInversion($\mathbb{F}_{p^2} \{x_0, x_1, ..., x_n - 1\}$)

```
1: for i = 0..(n-1) do
          den_i \leftarrow (x_i)_a^2 + (x_i)_b^2 \pmod{p}
 a_0 \leftarrow den_0
 4: for i = 1..(n-1) do
          a_i \leftarrow a_{i-1} \cdot den_i \pmod{p}
 6: inv \leftarrow a_{n-1}^{-1} \pmod{p}
 7: for i = n-1...1 do
          a_i \leftarrow inv \cdot dest_{i-1} \pmod{p}
 8:
          inv \leftarrow inv \cdot den_i \pmod{p}
 9:
10: a_0 \leftarrow a_{inv}
11: for i = 0..(n-1) do
          (xinv_i)_a \leftarrow a_i \cdot (x_i)_a \pmod{p}
12:
          (xinv_i)_b \leftarrow a_i \cdot -(x_i)_b \pmod{p}
13:
          x_i^{-1} \leftarrow \{(xinv_i)_a, (xinv_i)_b\}
14:
15: return \{x_0^{-1}, x_1^{-1}, ..., x_{n-1}^{-1}\}
```

- $2n \mathbb{F}_p$ squarings
- $n \mathbb{F}_p$ additions
- 1 \mathbb{F}_p inversion
- 3(n-1) \mathbb{F}_p multiplications
- $2n \mathbb{F}_p$ multiplications

And so, with C measuring the performance of **PartialBatchedInversion**, we have

$$C = 2n\mathbf{s} + n\mathbf{a} + \mathbf{i} + 3(n-1)\mathbf{m} + 2n\mathbf{m}$$

We can further simplify C if we presume that the execution time of squaring is roughly the same as multiplication. Additionally, we can simplify 3(n-1) to 3n in the spirit of complexity theory. With these simplifications we arrive at

$$C \approx 7n\mathbf{m} + n\mathbf{a} + \mathbf{i}$$

Applying the same simplifying assumptions to C_0 , we arrive at

$$C_0 \approx 3n\bar{\mathbf{m}} + 5\mathbf{m} + \mathbf{a} + \mathbf{i}$$

We note here that an \mathbb{F}_{p^2} multiplication $(\bar{\mathbf{m}})$ is performed simply by means of $4 \mathbb{F}_p$ multiplications (again, recall the multiplication of complex numbers). So we have $\bar{\mathbf{m}} = 4\mathbf{m}$, and can further simplify C_0 :

$$C_0 \approx (12n+5)\mathbf{m} + \mathbf{a} + \mathbf{i}$$

Finally we've simplified C and C_0 to forms that are more easily compared. Lets us turn our attention to the proposition that C runs in fewer operations than C_0 :

$$C < C_0$$

$$7n\mathbf{m} + n\mathbf{a} + \mathbf{i} < (12n + 5)\mathbf{m} + \mathbf{a} + \mathbf{i}$$

Simplifying slightly, we need now to resolve

$$7n\mathbf{m} + n\mathbf{a} < (12n + 5)\mathbf{m} + \mathbf{a}$$

 $n\mathbf{a} - \mathbf{a} < (12n + 5)\mathbf{m} - 7n\mathbf{m}$
 $n\mathbf{a} - \mathbf{a} < 5n\mathbf{m} + 5\mathbf{m}$
 $(n - 1)\mathbf{a} < (5n + 1)\mathbf{m}$

It appears now that in order for **PartialBatchedInv**₀ to be computationally favourable over **PartialBatchedInv**, the execution time for one \mathbb{F}_p addition would need to be larger than at least 5 times that of one \mathbb{F}_p multiplication.

Though it seems trivially true, we can verify this by measuring and comparing the execution times of the $\mathtt{SIDH}_{\mathtt{C}}$ addition and multiplication functions we will be using for our implementation.

When doing so (using the arithmetic test cases included in arith_tests.c by Microsoft Research) we arrive at the measurements outlined in table 3.1.3.

Operation	SIDH _C function	performance in clock cycles
\mathbb{F}_p addition	fpadd751	206
\mathbb{F}_p multiplication	fpmult751_mont	1,009

If we query for the performance of other operations (including \mathbb{F}_{p^2} arithmetic) we can estimate to what degree roughly **PartialBatchedInv** outperforms **PartialBatchedInv**₀. We can also measure to what degree we can expect that it will outperform an unbatched implementation of n-many inversions.

Operation	SIDH _C function	performance in clock cycles
\mathbb{F}_p inversion	fpinv751_mont	826,228
\mathbb{F}_{p^2} addition	fp2add751	172
\mathbb{F}_{p^2} multiplication	fp2mult751_mont	2,793
\mathbb{F}_{p^2} inversion	fp2inv751_mont	829,786

All of these results are computed as the average over 100 distinct applications. Furthermore, because they are measured in clock cycles, they are independent of any CPU clock rate. Because of this they are indicative of the complexity of each operation (or rather, the complexity of these implementations,) opposed to the performance of these operations on any one particular machine.

We conclude this section by using these results, along with the operation counts of each procedure, to compare the expected performances of **PartialBatchedInv**, **PartialBatchedInv**₀, and unbatched inversion. These results are shown in Table 3.1.3. For these estimations we have set the number of elements (n) equal to 248. This closely reflects the setting in which **PartialBatchedInv** will be implemented in SIDH_C, as will be discussed in the following section.

PartialBatchedInv₀:

If we substitute the performance variables in C_0 with the corresponding results from the tables above, we have:

$$C_0 \approx (12n+5)\mathbf{m} + \mathbf{a} + \mathbf{i}$$

 $C_0 \approx (12n+5)1,009 + 206 + 826,228$
 $C_0 \approx 12,108n + 831,479$

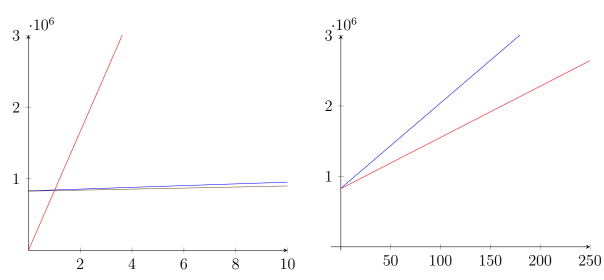
unbatched \mathbb{F}_{p^2} inversions:

The performance of *n*-many unbatched \mathbb{F}_{p^2} inversions can be modelled plainly by $n\bar{\mathbf{i}}$. The cost of *n* unbatched inversions is therefore 829, 786*n*.

PartialBatchedInv:

$$C \approx 7n\mathbf{m} + n\mathbf{a} + \mathbf{i}$$

 $C \approx 7,269n + 826,228$



Procedure	operation count	expected performance in clock cycles
PartialBatchedInv	fpinv751_mont	826,228
PartialBatchedInv ₀ 248 unbatched \mathbb{F}_{p^2} inversions	fp2add751 fp2inv751_mont	172 829,786

3.2 Implementation Details

We will now take the work of the previous subsection and explain in detail how it can be applied to the Yoo et al. signature layer of SIDH_C. We will begin with a high-level overview of the signature layer architecture. This mapping will allow efficient highlighting of the potential bottlenecks in the protocols performance. In this search for performance bottlenecks, we will specifically seek out execution paths in the codebase where-in batching inversions could offer a performance increase. Additionally, will discuss properties of

the signature scheme that can be leveraged to optimize the performance increases offered by **PartialBatchedInv**.

After this breif mapping of the signature layer, we move into the lower-level details of our procedures implementation. We transcribe **PartialBatchedInversion** to its C variant, pb_inv, which is defined almost entirely by means of the SIDH_C API. We will discuss some design specifics of pb_inv, and look breifly at the security of the function with respect to the signature scheme.

We round off this subsection with a discussion of other possible ways in which pb_inv (and operation batching in general) can offer performance improvements to the Yoo et al. signature scheme. This discussion is continued in the final Chapter of this dissertation, in Section 5.2.

3.2.1 Parallelizing Signatures

Recall Figure 2.6 which details the abstraction levels of the SIDH protocols as they relate to the modules of SIDH_c. We can further expand on this to figure to illustrate how the Yoo et al. signature layer interoperates with the original SIDH_c codebase. See Figure 3.1 - "SIDH_signature.c" signifies the code module added by Yoo et al., which implements the

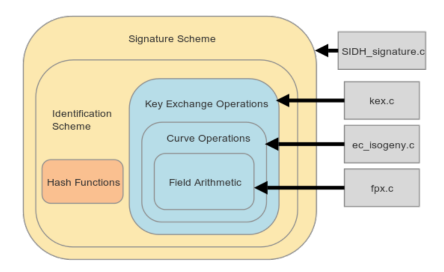


Figure 3.1: Relationship between SIDH based signatures & the Yoo et al. fork of the SIDH C library

Because every 2λ iteration of the **Sign** and **Verify** procedures are entirely independent of each other, these functions present themselves as embarrassingly parallel.²

Recall from section 2.6.4 the table of functions that can be found in SIDH_signature. isogey_sign acts as the entry point for Sign and spawns a POSIX thread for every instance of the for-loop; each one calling sign_thread which performs Bob's interaction with Randall. Verification proceeds analogously; isogeny_verify is executed and spawns a POSIX threads executing verify_thread until all 2λ iterations are complete.

 $^{^{2}}$ in the field of high performance computing, a problem that is trivially parallizable is often referred to as embarrassingly parallizable.

This parallelization of the signature scheme was the approach taken by Yoo et al. in their original implementation. It also lends itself rather nicely

inv_4_way .

```
void inv_4_way(f2elm_t z1, f2elm_t z2, f2elm_t z3, f2elm_t z4) {
1
2
        f2elm_t t0, t1, t2;
3
        int tempCnt;
4
                                                              // t0 = z1*z2
5
        fp2mul751_mont(z1, z2, t0);
                                                              // t1 = z3*z4
6
        fp2mul751_mont(z3, z4, t1);
                                                              // t2 = z1*z2*z3*z4
        fp2mul751_mont(t0, t1, t2);
7
8
        fp2inv751\_mont(t2);
                                                              // t2 = 1/(z1*z2*z3*z4)
9
        fp2mul751\_mont(t0, t2, t0);
                                                              // t0 = 1/(z3*z4)
10
                                                              // t1 = 1/(z1*z2)
        fp2mul751_mont(t1, t2, t1);
                                                              // t2 = 1/z4
11
        fp2mul751_mont(t0, z3, t2);
                                                             // z3 = 1/z3
12
        fp2mul751_mont(t0, z4, z3);
                                                             // z_4 = 1/z_4
13
        fp2copy751(t2, z4);
                                                              // t2 = 1/z2
14
        fp2mul751_mont(z1, t1, t2);
                                                             // z1 = 1/z1
// z2 = 1/z2
        fp2mul751_mont(z2, t1, z1);
15
16
        fp2copy751(t2, z2);
17
   j_inv.
```

```
void j_inv(const f2elm_t A, const f2elm_t C, f2elm_t jinv) {
2
        f2elm_t t t0, t1;
3
        fp2sqr751_mont(A, jinv);
                                                                  // jinv = A^2
4
                                                                    t1 = C^2
5
        fp2sqr751\_mont(C, t1);
6
        fp2add751(t1, t1, t0);
                                                                  // t0 = t1+t1
7
        fp2sub751(jinv, t0, t0);
                                                                  // t\theta = jinv - t\theta
8
        fp2sub751(t0, t1, t0);
                                                                  // t0 = t0-t1
9
        fp2sub751(t0, t1, jinv);
                                                                  // jinv = t0-t1
10
        fp2sqr751_mont(t1, t1);
                                                                  // t1 = t1^2
                                                                  // jinv = jinv*t1
11
        fp2mul751_mont(jinv, t1, jinv);
                                                                  // t\theta = t\theta + t\theta
        fp2add751(t0, t0, t0);
12
        fp2add751(t0, t0, t0);
                                                                  // t\theta = t\theta + t\theta
13
14
        fp2sqr751_mont(t0, t1);
                                                                  // t1 = t0^2
        fp2mul751_mont(t0, t1, t0);
                                                                  // t0 = t0*t1
15
                                                                  //t0 = t0+t0
16
        fp2add751(t0, t0, t0);
17
        fp2add751(t0, t0, t0);
                                                                  // t0 = t0+t0
18
        fp2inv751_mont(jinv);
                                                                   // jinv = 1/jinv
19
        fp2mul751_mont(jinv, t0, jinv);
                                                                  // jinv = t0*jinv
20
```

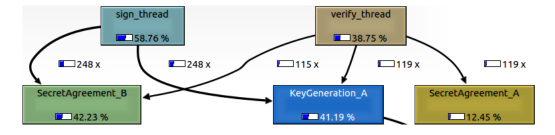


Figure 3.2: ¡Caption here;

sign_thread verify_thread

SecretAgreement_B KeyGeneration_A SecretAgreement_B

Because the work of Yoo et al. was built on top of the original Microsoft SIDH library, all underlying field arithmetic (and as such, pointwise arithmetic) is performed in projective space using Montgomery representation. As was mentioned in the previous chapter, doing such allows us to avoid a great deal of field element inversions. We simply convert to Montgomery representation at the beginning of heavy arithmetic, perform the desired operations, and then convert back to standard representation when complete. Throughout the codebase, (most noticeably in kex.c) we can see an analogous approach taken in for pointwise arithmetic, by first converting to a projective space representation, performing operations, and then converting back to affine coordinates.

The downside of this (for our work) is that the number opportunities for implementing the batched inversion algorithm becomes greatly limited.

3.2.2 Implementation & Design Decisions

With Algorithm ?? we provide an explicit C definition for the function pb_inv. For descriptions of the functions called in this procedure, the reader can refer to section 2.6.3. For explicit definitions of some of these functions, the reader can refer to Appendix A.

Security Considerations. Recall the notion of a general side-channel attack: A side-channel attack is performed when an unauthorized individual is able to acquire information by measuring properties of the physical implementation of the system at hand. This can be done by analyzing the power consumption, timing properties, or electromagnetic leaks of a CPU while it operates on (or generates) confidential information.

In the context of information security, algorithms for performing operations over mathematical objects can be said to fall under one of two categories: *constant time* and *non-constant time* algorithms. Constant time algorithms are designed to protect confidential information from side-channel attacks, but come at the cost of computational efficiency.

In the SIDH_C library, there are two distinct functions for computing field element inversions: fp2inv751_mont and fp2inv751_mont_bingcd. fp2inv751_mont_bingcd performs inversion by means of the binary GCD (greatest common denominator) algorithm, and is a non-constant time implementation. fp2inv751_mont is a constant time implementation, and as such runs slower than fp2inv751_mont_bingcd in nearly all cases, but protects against timing based side-channel attacks. They perform comparatively as such:

Procedure	Performance in clock cyckes	System A With Batching
fp2inv751_mont	68,881,331	68,881,331
fp2inv751_mont_bingcd	15,744,477,032	15,565,738,003

Take for example some private data c being manipulated or operated on by some algorithm **A**. In order to be entirely certain that c in $\mathbf{P}(c)$ is not vulnerable to any

```
void pb_inv (const f2elm_t* vec, f2elm_t* dest, const int n) {
1
2
        felm_t t0[n];
                            //a portion of vec elements
3
        felm_t t1[n];
                            //b portion of vec elements
4
        felm_t den[n];
                            //denominator of vec elements
5
        felm_t a[n];
6
7
        // upward-percolation phase -
8
9
        for (int i = 0; i < n; i++) {
10
            //compute \ denp[i] = a[i][0]^2 + a[i][1]^2
             fpsqr751_mont((vec[i])[0], t0[i]);
11
12
             fpsqr751_mont((vec[i])[1], t1[i]);
            fpadd751(t0[i], t1[i], den[i]);
13
        }
14
15
        fpcopy751\left(\,den\left[\,0\,\right]\,,\;\;a\left[\,0\,\right]\,\right);
16
17
        for (int i = 1; i < n; i++) {
18
            fpmul751_mont(a[i-1], den[i], a[i]);
19
20
21
        // inversion phase -
22
23
        felm_t a_inv;
24
        fpcopy751(a[n-1], a_inv);
25
        fpinv751_mont_bingcd(a_inv);
26
        // downward percolation phase
27
                                                                              -//
28
29
        for (int i = n-1; i >= 1; i--) {
             fpmul751_mont(a[i-1], a_inv, a[i]);
30
31
            fpmul751_mont(a_inv, den[i], a_inv);
32
        }
33
34
        fpcopy751 (a_inv , a [0]);
35
36
        for (int i = 0; i < n; i++) {
             //convert back to extension field
37
            fpmul751_mont(a[i], vec[i][0], dest[i][0]);
38
39
            fpneg751 ((vec[i])[1]);
40
            fpmul751_mont(a[i], vec[i][1], dest[i][1]);
41
42
```

Figure 3.3: pb_inv

imagineable side-channel attack it must be the case that the structure of \mathbf{P} does not in anyway depend on the information stored in c.

Let us look more closely at the elements that are inverted by pb_inv. In KeyGeneration_A the elements being passed to pb_inv are simply the constituents of Randall's public key for every iteration of the signing procedure. Because the data being operated on is publicly available information, we needn't consider if it is vulnerable to side-channel analysis.

In SecretAgreement_A, pb_inv is called by j_inv, which produces the j-invariant of a particular curve. When SecretAgreement_A is used in the context of SIDH key exchange, j_inv is used to compute the shared secret between party members A and B, and so would need to be protected against side-channel attacks. This is not the case in the context of signatures, however. Recall the structure of SIDH signatures: every signature includes, for every 2λ iterations, the commitment E_1 , which is precisely the shared secret between the signer and Randall. And so, every use of pb_inv in the context of signatures does not require extra precautions to side-channel analysis. The same is true for SecretAgreement_B.

While it is not the case for our implementations that the data on which pb_inv operates would find itself the subject of side-channel analysis, it is not difficult to imagine a scenario where this might be the case.

3.3 Results

Modulus Size	Regular Batch	Partial Batched Inversion	Unbatched
32 Inversion 64 Inversion 128 Inversion 256 Inversion 512 Inversion	0.033685 0.00033685 0.00033685 0.00033685 0.00033685	0.00080204 0.00080204 0.00080204 0.00080204 0.00080204	
1024 Inversion 2048 Inversion	0.00033685 0.00033685	0.00080204 0.00080204	

Modulus Size	Regular Batch	Partial Batched Inversion	Unbatched
32	0.00033685	0.00080204	
64	0.00033685	0.00080204	
128	0.00033685	0.00080204	
256	0.00033685	0.00080204	
512	0.00033685	0.00080204	
1024	0.00033685	0.00080204	
2048	0.00033685	0.00080204	

Modulus Size	Regular Batch	Partial Batched Inversion	Unbatched
32	0.00033685	0.00080204	
64	0.00033685	0.00080204	
128	0.00033685	0.00080204	
256	0.00033685	0.00080204	
512	0.00033685	0.00080204	
1024	0.00033685	0.00080204	
2048	0.00033685	0.00080204	

Two different machines were used for benchmarking. System A denotes a single-core, 1.70 GHz Intel Celeron CPU. System B denotes a quad-core, 3.1 GHz AMD A8-7600.

The two figures below provide benchmarks for KeyGen, Sign, and Verify procedures with both batched partial inversion implemented (in the previously mentioned locations) and not implemented. All benchmarks are averages computed from 100 randomized sample runs. All results are measured in clock cycles.

Procedure	System A Without Batching	System A With Batching
KeyGen Signature Sign	68,881,331 15,744,477,032	68,881,331 15,565,738,003
Signature Verify	11,183,112,648	10,800,158,871
Procedure	System B Without Batching	System B With Batching
KeyGen	84,499,270	84,499,270
Signature Sign	10,227,466,210	10,134,441,024
Signature Verify	7,268,804,442	7,106,663,106

System A: With inversion batching turned on we notice a 1.1% performance increase for key signing and a 3.5% performance increase for key verification.

System B: With inversion batching turned on we a observe a 0.9% performance increase for key signing and a 2.3% performance increase for key verification.

3.3.1 Analysis

It should first be noted that, because our benchmarks are measured in terms of clock cycles, the difference between our two system clock speeds should be essentially ineffective.

In the following table, "Batched Inversion" signifies running pb_{inv} on 248 \mathbb{F}_{p^2} elements.

Procedure	Performance
Batched Inversion \mathbb{F}_{p^2} Montgomery Inversion	1721718 874178

The following are all measured in clock cycles, as the computed average of 1000 distinct executions:

Modulus Size	Multiplication	Inversion Time
32	0.00033685	0.00080204
64	0.00033685	0.00080204
128	0.00033685	0.00080204
256	0.00033685	0.00080204
512	0.00033685	0.00080204
1024	0.00033685	0.00080204
2048	0.00033685	0.00080204

Do performance increases observed make sense?

3.3.2 Remaining Opportunities

There are two functions called in the isogeny signature system that perform a \mathbb{F}_{p^2} inversion: j_{inv} and inv_4_way . These functions are called once in SecretAgreement and KeyGeneration operations respectively. SecretAgreement and KeyGeneration are in turn called from each signing and verification thread.

This means that in the signing procedure there are 2 opportunities for implementing batched partial-inversion with a batch size of 248 elements. In the verify procedure, however, there are 3 opportunities for implementing batched inversion with a batch size of roughly 124 elements.

Another avenue for implementation of this procedure, much in line with Fiat's original RSA batch, lies in . We discuss this approach in greater detail in section 5.2.

Chapter 4

Compressing Signatures

Costello et al. showed in [CJL⁺17] that

4.1 SIDH Key Compression Background

We discussed rejection sampling A values from signature public keys until we found an A that was also the x-coord of a point. After some simple analysis, however, we found that it was extremely unlikely for A to be a point on the curve.

- 4.1.1 Motivation & Overview
- 4.1.2 Construction of Bases
- 4.1.3 Pohlig-Hellman
- 4.1.4 Decompression
- 4.2 Implementation Details
- 4.2.1 Tailoring Compression for Signatures
- **4.2.2** Decompressing $\psi(S)$
- 4.3 Results
- 4.4 Analysis

Chapter 5

Discussion & Conclusion

- 5.1 Results & Comparisons
- 5.2 Additional Opportunities for Batching
- 5.3 Future Work

¡Conclusion here;

Appendices

$\begin{array}{c} \mathbf{Appendix} \ \mathbf{A} \\ \\ \mathtt{SIDH}_{\mathbb{C}} \ \mathbf{Functions} \end{array}$

here's an appendix

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