ANALYSIS OF PSLQ, AN INTEGER RELATION FINDING ALGORITHM

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ABSTRACT. Let \mathbb{K} be either the real, complex, or quaternion number system and let $\mathbb{O}(\mathbb{K})$ be the corresponding integers. Let $x = (x_1, \ldots, x_n)$ be a vector in \mathbb{K}^n . The vector x has an integer relation if there exists a vector $m = (m_1, \ldots, m_n) \in \mathbb{O}(\mathbb{K})^n$, $m \neq 0$, such that $m_1x_1 + m_2x_2 + \ldots + m_nx_n = 0$. In this paper we define the parameterized integer relation construction algorithm PSLQ(τ), where the parameter τ can be freely chosen in a certain interval.

Beginning with an arbitrary vector $x = (x_1, \ldots, x_n) \in \mathbb{K}^n$, iterations of $PSLQ(\tau)$ will produce lower bounds on the norm of any possible relation for x. Thus $PSLQ(\tau)$ can be used to prove that there are no relations for x of norm less than a given size. Let M_x be the smallest norm of any relation for x. For the real and complex case and each fixed parameter τ in a certain interval, we prove that $PSLQ(\tau)$ constructs a relation in less than $O(n^3 + n^2 \log M_x)$ iterations.

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1. INTRODUCTION

Let \mathbb{K} be either the real, complex or quaternion number system and let $\mathbb{O}(\mathbb{K})$ be the corresponding system of integers (i.e., ordinary integers, Gaussian integers, or Hamiltonian integers, respectively). Let $x = (x_1, \ldots, x_n)$ be a vector in \mathbb{K}^n . The vector x has an *integer relation* if there exists a vector $m = (m_1, \ldots, m_n) \in \mathbb{O}(\mathbb{K})^n, m \neq 0$, such that $m_1 x_1 + m_2 x_2 + \ldots + m_n x_n = 0$.

In this paper we define the parameterized integer relation construction algorithm $PSLQ(\tau)$, which, compared with other integer relation algorithms in the literature, features superior performance and excellent numerical stability. The parameter τ can be freely chosen in the interval $1 < \tau < \rho$, where ρ is 2 or $\sqrt{2}$ depending on whether K is the reals or complexes, respectively; if K is the quaternions take τ and ρ to be 1. We analyze $PSLQ(\tau)$ for these three number systems. We describe in detail some efficient Fortran multiprecision computer implementations of $PSLQ(\tau)$.

Beginning with an arbitrary vector $x = (x_1, \ldots, x_n) \in \mathbb{K}^n$, a finite number of iterations of $PSLQ(\tau)$ will produce lower bounds on the (Frobenius) norm of any possible relation for x. The computation of such a lower bound constitutes a proof that x has no integer relations whatsoever of norm less than this lower bound. Any finite computation done with PSLQ or any other presently known relation finding algorithm can only prove that no small relation exists. Such an algorithm can construct an alleged relation based on inputs given to finite precision, but the proof that this alleged relation is a true relation for the real numbers is a separate matter.

Let M_x be the smallest norm of a relation for x. Define γ by the equation $\tau = 1/\sqrt{1/\rho^2 + 1/\gamma^2}$, where $\rho = 2$ for the real number field and $\rho = \sqrt{2}$ for the complex number field, and τ as above. For each fixed parameter τ in the interval $1 < \tau < \rho$, we prove in the real and complex case that $\text{PSLQ}(\tau)$ constructs a relation in less than $\binom{n}{2} \log_{\tau} (\gamma^{n-1} M_x)$ iterations. This shows that $\text{PSLQ}(\tau)$ is 'polynomial time' in the dimension and the number of bits of a smallest integer relation. Different τ or γ choices lead to different time and space requirements for the algorithm.

For dimension n = 2 we prove that $PSLQ(\tau)$ will construct a relation of smallest norm M_x . We give examples in dimension n = 3, for some τ , for which $PSLQ(\tau)$ does not construct a relation of smallest norm M_x . However for any dimension $n \ge 2$, we do prove that any relation constructed by $PSLQ(\tau)$ has norm less than or equal to $\gamma^{n-2}M_x$.

The 'polynomial time' and 'small norm' proofs given here are straightforward generalizations to the parameter τ and to the complex numbers of the original 'polynomial time' proofs which appear in Lagarias et al, [23].

We show, however, that the algorithm of [23] is distinct from any of these $PSLQ(\tau)$ algorithms.

 $PSLQ(\tau)$ was introduced by the authors [2] in 1991. PS refers to partial sums of squares, LQ to a lower trapezoidal orthogonal decomposition, and (τ) is a parameter defined as above. Since $PSLQ(\tau)$ was introduced it has been used to discover numerous previously unknown identities among real numbers. One example is

$$\sum_{k=1}^{\infty} \left(1 - \frac{1}{2} + \dots + \frac{(-1)^{k+1}}{k} \right)^2 (k+1)^{-3}$$
$$= 4L_5(1/2) - \frac{1}{30} \ln^5(2) - \frac{17}{32}\zeta(5) - \frac{11}{720}\pi^4 \ln(2)$$
$$+ \frac{7}{4}\zeta(3) \ln^2(2) + \frac{1}{18}\pi^2 \ln^3(2) - \frac{3}{24}\pi^2\zeta(3),$$

where $L_n(x)$ denotes the polylogarithm function $\sum_k x^k k^{-n}$. See [3] for details. Another example is the following formula for π :

$$\pi = \sum_{i=0}^{\infty} \frac{1}{16^i} \left(\frac{4}{8i+1} - \frac{2}{8i+4} - \frac{1}{8i+5} - \frac{1}{8i+6} \right)$$

This remarkable series permits one to rapidly compute individual digits from the hexadecimal expansion of π . See [4] for details. It was found by applying PSLQ(τ) to the vector $X = (X_1, X_2, \dots, X_8, \pi)$ where $X_j = \sum_{k>0} 1/(16^k(8k+j))$. The smallest relation known,

$$(4, 0, 0, -2, -1, -1, 0, 0, -1)$$

yields the above 'base 16' formula for π . A next smallest relation known,

$$(0, 8, 4, 4, 0, 0, -1, 0, -2),$$

was subsequently discovered by Ferguson and this relation yields a similar 'base 16' formula for π . Together these two integral lattice relation vectors generate a two-dimensional lattice of relations of this 'base 16' type. It is conjectured there are no further such relations outside this lattice. Note that

$$(-8, 8, 4, 8, 2, 2, -1, 0, 0)$$

is in this lattice, so evidently X_7 is integrally dependent upon X_1, \ldots, X_6 . 3 Of course, a numerical discovery of a relation using $PSLQ(\tau)$ does not constitute a rigorous proof of the relation. However, in the wake of this numerical evidence, proofs have subsequently been found for many of these relations, including the above formula for π . See [3] and [4] for details.

In the theoretical proofs in Section 2, 3, 4, and 5, we will assume exact arithmetic over the real numbers augmented by comparisons over the reals and the nearest integer function.

2. Lower Bounds on Integer Relations

If K is the complex number field, then z^* denotes the complex conjugate of z, i.e. if z = x + iy, then $z^* = x - iy$. $|\cdot|$ denotes the complex absolute value, i.e. $|z|^2 = z^*z = zz^* = x^2 + y^2$. If A is a matrix or vector, then A^* is the conjugate transpose of A. A unit in the complex number field is any element z such that |z| = 1. For real z, the conjugate operation is null, and z is the usual absolute value.

Similarly, if K is the quaternion number system, then z^* denotes the quaternion conjugate of z, i.e. if z = x+yi+uj+vk, then $z^* = x-yi-uj-vk$. The quaternion absolute value or norm is similarly defined, so that $|z|^2 = zz^* = z^*z = x^2 + y^2 + u^2 + v^2$. Units and conjugates of matrices are defined analogously.

If K is any of the above three number systems, two vectors $x, y \in \mathbb{K}^n$ are said to be orthogonal if $xy^* = 0$. Let $|A| = (\operatorname{tr}(A^*A))^{1/2}$ denote the Frobenius norm of the matrix A, i.e., $|A| = (\sum a_{i,j}^* a_{i,j})^{1/2}$. An $n \times n$ matrix A is unitary if $A^*A = AA^* = I_n$. $U(n, \mathbb{K})$ denotes the group of unitary matrices over K. An $n \times n$ matrix A is unimodular if det A is a unit. $GL(n, \mathbb{O}(\mathbb{K}))$ is the group of unimodular matrices with entries in the integers $\mathbb{O}(\mathbb{K})$.

Definition 1: (M_x) . Assume $x = (x_1, \ldots, x_n) \in \mathbb{K}^n$ has norm |x| = 1. Define x^{\perp} to be the set of all vectors in \mathbb{K}^n orthogonal to x. Let $\mathbb{O}(\mathbb{K})^n \cap x^{\perp}$ be the discrete lattice of integral relations for x. Define $M_x > 0$ to be the smallest norm of any relation for x in this lattice.

Definition 2: (H_x) . Assume $x = (x_1, \ldots, x_n) \in \mathbb{K}^n$ has norm |x| = 1. Furthermore, suppose that no coordinate entry of x is zero, i.e., $x_j \neq 0$ for $1 \leq j \leq n$ (otherwise x has an immediate and obvious integral relation). For $1 \leq j \leq n$ define the partial sums

$$s_j^2 = \sum_{\substack{j \le k \le n \\ 4}} x_k x_k^\star.$$

Given such a unit vector x define the $n \times (n-1)$ lower trapezoidal matrix $H_x = (h_{i,j})$ by

$$h_{i,j} = \begin{cases} 0 & \text{if } 1 \le i < j \le n-1 \\ s_{i+1}/s_i & \text{if } 1 \le i = j \le n-1 \\ -x_i^* x_j/(s_j s_{j+1}) & \text{if } 1 \le j < i \le n. \end{cases}$$

Note that $h_{i,j}$ is scale invariant.

Lemma 1. Let H_x be the lower trapezoidal matrix defined above. Then (i) $H_x^*H_x = I_{n-1}$, i.e., the columns of H_x are orthogonal, (ii) $|H_x| = \sqrt{n-1}$, (iii) $xH_x = 0$.

Proof. The columns can be proven orthogonal by considering the cases i = j and i < j separately. When i = j the inner product is

$$\frac{s_{i+1}^2}{s_i^2} + \sum_{i < k \le n} \frac{x_i x_i^* x_k x_k^*}{s_i^2 s_{i+1}^2} = \frac{s_{i+1}^2}{s_i^2} + \frac{x_i x_i^*}{s_i^2 s_{i+1}^2} \sum_{i < k \le n} x_k x_k^*$$
$$= \frac{s_i^2 - x_i x_i^*}{s_i^2} + \frac{x_i x_i^*}{s_i^2} = 1.$$

When i < j the inner product is

$$-\frac{s_{j+1}x_i^* x_j}{s_j s_i s_{i+1}} + \sum_{j < k \le n} \frac{x_i^* x_j x_k x_k^*}{s_i s_{i+1} s_j s_{j+1}}$$
$$= -\frac{s_{j+1}x_i^* x_j}{s_j s_i s_{i+1}} + \frac{x_i^* x_j}{s_i s_{i+1} s_j s_{j+1}} \sum_{j < k \le n} x_k x_k^* = 0.$$

Item (i) shows that $H_x^* H_x = I_{n-1}$ which has trace n-1 so $|H_x| = \sqrt{n-1}$. To prove (iii), fix $1 \le j \le n-1$, then

$$\sum_{1 \le k \le n} x_k h_{k,j} = \frac{x_j s_{j+1}}{s_j} - \sum_{j < k \le n} \frac{x_k x_k^* x_j}{s_j s_{j+1}} = \frac{x_j s_{j+1}^2}{s_j} - \frac{x_j s_{j+1}^2}{s_j s_{j+1}} = 0.$$

Lemma 2. For a unit vector $x \in \mathbb{K}^n$ define $P_x = H_x H_x^*$. Then P_x satisfies: (i) $P_x^* = P_x$, (ii) $P_x = I_n - x^*x$, (iii) $P_x^2 = P_x$, (iv) $|P_x| = \sqrt{n-1}$, (v) $P_x z^* = z^*$ for any $z \in x^{\perp}$ (vi) $P_x m^* = m^*$ for any relation $m \in \mathbb{O}(\mathbb{K})^n$ for x.

Proof. Item (i) follows from $H_x H_x^* = (H_x H_x^*)^*$. To prove (ii) note that from Lemma 1 (iii), H_x is an $n \times (n-1)$ rank n-1 matrix whose columns transposed form an orthonormal basis for x^{\perp} . Defining $U = (H_x | x^*)$, an $n \times n$ unitary matrix, we have $UU^* = H_x H_x^* + x^* x = I_n$. To prove (iii) note that

$$P_x^2 = (I_n - x^* x)^2 = I_n^2 - 2I_n x^* x + x^* (xx^*) x = P_x.$$

To prove (iv) note that $|P_x|^2 = \operatorname{tr}(P_x^*P_x) = \operatorname{tr}P_x = \operatorname{tr}H_x^*H_x = n-1$. Item (vi) follows from (v) which follows from (ii) and the associativity $(x^*x)z^* = x^*(xz^*)$. \Box

Theorem 1. Let $x \neq 0 \in \mathbb{K}^n$. Suppose that for any relation m of x and for any matrix $A \in GL(n, \mathbb{O}(\mathbb{K}))$ there exists a unitary matrix $Q \in U(n-1)$ such that $H = AH_xQ$ is lower trapezoidal and all of the diagonal elements of H, $h_{j,j} \neq 0$. Then

$$\frac{1}{\max_{1 \le j \le n-1} |h_{j,j}|} = \min_{1 \le j \le n-1} \frac{1}{|h_{j,j}|} \le |m|$$

Proof. Let m be any relation for x. By the hypothesis, there exists a unitary matrix $Q \in U(n-1)$ such that $H = AH_xQ$ is lower trapezoidal (this is equivalent to QR factorization). There is an $n \times n - 1$ matrix T with diagonal ones and an $n - 1 \times n - 1$ diagonal matrix D where H = TD with diagonal entries $h_{j,j} \neq 0, 1 \leq j \leq n - 1$ from the hypothesis. On the other hand, $AP_x = HQ^*H_x^*$, from the definition of P_x in Lemma 2. The equation $AP_x = TDQ^*H_x^*$ gives a decomposition of AP_x into the product of a lower trapezoidal matrix T with diagonal 1's, an invertible diagonal matrix D with diagonal h's, and an $n - 1 \times n$ matrix $Q^*H_x^*$ with orthonormal rows since $Q^*H_x^*H_xQ = Q^*I_{n-1}Q = I_{n-1}$ by Lemma 1. So the norm of the j-th row of $DQ^*H_x^*$ is $|h_{j,j}|$.

From Lemma 2, part (vi), $m^* = P_x m^*$, so that $Am^* = AP_x m^*$. From the above decomposition of $AP_x = TDQ^*H_x^*$, we have $Am^* = AP_x m^* = \frac{1}{6}$

 $TD(Q^{\star}H_x^{\star})m^{\star}$. Let $Q_{H,j}$ be the *j*-th row of $Q^{\star}H_x^{\star}$ and let A_j be the *j*-th row of A. Then

$$A_j m^{\star} = h_{j,j} Q_{H,j} m^{\star} + \sum_{k < j} t_{j,k} h_{k,k} Q_{H,k} m^{\star}.$$

Since A is invertible, $Am^* \neq 0$. Let j be the least j for which $A_jm^* \neq 0$ so that $A_km^* = 0$ for k < j. Then the k < j rows of $TDQ^*H_x^*m^*$ are zero, and since T is lower trapezoidal by recursion, the k-th rows of $Q^*H_x^*m^*$ are also zero. With this least choice of j then $A_jm^* = h_{j,j}Q_jm^*$. Therefore, from $A \in GL(n, \mathbb{O}(\mathbb{K}))$,

$$1 \le |A_j m^*| \le |h_{j,j} Q_{H,j} m^*| \le |h_{j,j}| |m^*|,$$

because $Q_{H,i}$ is a unit vector. \Box

Comment on Theorem 1. Theorem 1 suggests a strategy to construct a relation finding algorithm: Find a way to reduce the norm of the matrix H_x by multiplication by some unimodular A on the left. The inequality of Theorem 1 offers an increasing lower bound on the size of any possible relation. Theorem 1 can be used with any algorithm that produces any $GL(n, \mathbb{O}(\mathbb{K}))$ matrices. Any $GL(n, \mathbb{O}(\mathbb{K}))$ matrix A whatsoever can be put into Theorem 1.

Definition 3: (Hermite reduction). Let H be a lower trapezoidal matrix, with $h_{i,j} = 0$ if j > i and $h_{j,j} \neq 0$. Define the matrix $D = (d_{i,j}) \in$ $GL(n, \mathbb{O}(K))$ recursively as follows. For fixed i, decrement j from n to 1, setting

$$d_{i,j} = \begin{cases} 0 & \text{if } i < j \\ 1 & \text{if } i = j \\ \operatorname{nint}((-\sum_{j < k \le i} d_{i,k}h_{k,j})/h_{j,j}) & \text{if } j < i, \end{cases}$$

We will say that DH is the *Hermite reduction* of H and we will say that D is the *reducing matrix* of H. The function nint denotes a nearest integer function, e.g., $nint(t) = \lfloor t + 1/2 \rfloor$. This definition of nint can be extended to each coordinate for complex or quaternion arguments.

Definition 4: (Modified Hermite reduction). With the same notation as in Definition 3, set $D = I_n$. For *i* from 2 to *n*, and for *j* from i - 1 to 1 (step -1), set $q = \text{nint}(h_{i,j}/h_{j,j})$; then for *k* from 1 to *j* replace $h_{i,k}$ by $h_{i,k} - qh_{j,k}$, and for *k* from 1 to *n* replace $d_{i,k}$ by $d_{i,k} - qd_{j,k}$.

Lemma 3. For a lower triangular matrix H with $h_{i,j} = 0$ if j > i and $h_{j,j} \neq 0$, Hermite reduction is equivalent to modified Hermite reduction.

Comment. This variation can be found in [8] and later in [28]. This recursion replaces the input H with DH while developing the left multiplying reduction matrix D.

Lemma 4. There exists a constant $\rho_{\mathbb{K}} = \rho \geq 1$, with the property that the entries of the Hermite reduced matrix $H' = (h'_{i,j}) = DH$ satisfy the inequality

$$|h'_{k,i}| \le |h'_{i,i}|/
ho = |h_{i,i}|/
ho$$

for all k > i. The constant $\rho = 2$ for the real case, $\rho = \sqrt{2}$ for the complex case, and $\rho = 1$ for the quaternion case.

Proof. This follows from the definitions of the nint function, Hermite reduction, and the fact that $|z - \operatorname{nint}(z)| \leq \sqrt{\dim_{\mathbb{R}} \mathbb{K}}/2$ for $z \in \mathbb{K}$. \Box

3. Statement of the Algorithm $PSLQ(\tau)$

Definition 5: (The parameters γ and τ). Fix the real number $\gamma > 2/\sqrt{3}$ or $\gamma > \sqrt{2}$ or $\gamma = \infty$ for the real, complex, and quaternion cases respectively. In terms of this γ , define the real number τ by

$$1/\tau^2 = 1/\rho^2 + 1/\gamma^2,$$

where ρ is defined as in Lemma 4. For the proof of Theorem 2, we will require that $1 < \tau$ and that $\tau \leq \rho$; clearly these conditions are satisfied in the real and complex cases. In the quaternion case $\tau = 1$ and $\rho = 1$.

For the proofs that follow assume \mathbb{K} is real or complex, not quaternion. Note however that the statement of the algorithm is valid for the quaternions.

Initial conditions: Given the input unit vector $x \in \mathbb{K}^n$, set $H = H_x$ where H_x is defined as above. Set the $n \times n$ matrices A and B to the identity I_n . Perform Hermite reduction on H, producing $D \in GL(n, \mathbb{O}(\mathbb{K}))$. Replace x by xD^{-1} , H by DH, A by DA, B by BD^{-1} .

One four-step iteration:

Step 1: Exchange

Let $H = (h_{i,j})$ where $h_{i,j}$ is the *i*-th row, *j*-th column entry of H. Let

$$\alpha = h_{r,r}, \quad \beta = h_{r+1,r}, \quad \lambda = h_{r+1,r+1}, \quad \delta = \sqrt{\beta\beta^{\star} + \lambda\lambda^{\star}}.$$

Choose an integer r such that $\gamma^r |h_{r,r}| \ge \gamma^i |h_{i,i}|$ for all $1 \le i \le n-1$. Define the permutation matrix R to be the identity matrix with the r and r+1rows exchanged. Replace x by xR, H by RH, A by RA, and B by BR.

Step 2: Corner

At this point the updated matrix H may not be lower trapezoidal since λ may not be zero. If r < n - 1 replace H by HQ where Q is the unitary $n - 1 \times n - 1$ matrix $Q = (q_{i,j}) \in U(n - 1, \mathbb{K})$ defined by

$$q_{i,j} = \begin{cases} \beta^{\star}/\delta & \text{if } i = r, j = r \\ -\lambda/\delta & \text{if } i = r, j = r+1 \\ \lambda^{\star}/\delta & \text{if } i = r+1, j = r \\ \beta/\delta & \text{if } i = r+1, j = r+1 \\ 1 & \text{if } i = j \neq r \text{ or } i = j \neq r+1 \\ 0 & \text{otherwise.} \end{cases}$$

where the $\alpha, \beta, \lambda, \delta$ are defined in Step 1. If r = n - 1 then H is unchanged.

Step 3: Reduction

Perform Hermite reduction on H, producing $D \in GL(n, \mathbb{O}(\mathbb{K}))$. Replace x by xD^{-1} , H by DH, A by DA, B by BD^{-1} .

Step 4: Termination

Terminate the algorithm if $x_j = 0$ for some $1 \le j \le n$ or if $h_{i,i} = 0$ for some $1 \le i \le n-1$.

4. Number of Iterations of $PSLQ(\tau)$

Let H(k) = H, A, and $B = A^{-1}$ be the result after exactly k iterations of PSLQ. Let $\alpha = h_{r,r}(k)$ and $\beta = h_{r+1,r}(k)$. These definitions of α and β are consonant with those of Step 2. Because H is Hermite reduced in Step 3, from Lemma 4, $|\beta| < |\alpha|/\rho$. For r < n-1 set $\lambda = h_{r+1,r+1}(k)$ and define t by $t = \sqrt{\beta\beta^* + \lambda\lambda^*}/|\alpha|$. From this definition of t we have

$$|\lambda| \le |\alpha|t.$$

From the Step 1 Exchange, $0 \leq |\lambda| \leq |\alpha|/\gamma$. It follows that

$$t = \sqrt{\beta\beta^* + \lambda\lambda^*}/|\alpha| \le \sqrt{1/\rho^2 + 1/\gamma^2} = \tau,$$

as in Definition 5. For this proof we will require that $t < 1 < \tau$, clearly satisfied in the real and complex cases.

Lemma 5. If $h_{j,j}(k) = 0$ for some $1 \le j \le n-1$ and no smaller k, then j = n-1 and a relation for x must appear as a column of the matrix B.

Proof. (Alyson Reeves) First we show that $h_{j,j} = 0$ implies that j = n - 1. Consider the matrix H(k-1), the end result of the k-1-th iteration. By the hypothesis on k we know that no diagonal elements in H(k-1) are zero. In particular, for the r about to be chosen in Step 1 of the k-th iteration, we know that $h_{r,r}(k-1) \neq 0$ and that $h_{r+1,r+1}(k-1) \neq 0$. Now, suppose the r chosen in Step 1 is not n-1. Let

$$\begin{pmatrix} \alpha & 0 \\ \beta & \lambda \end{pmatrix}$$

be the submatrix of H(k-1) consisting of the r and r+1 rows of columns r and r+1. After Step 1 has been performed this submatrix becomes

$$\begin{pmatrix} \beta & \lambda \\ \alpha & 0 \end{pmatrix}.$$

At Step 2, we post-multiply the matrix by the unitary sub-matrix of Q

$$\begin{pmatrix} \beta^{\star}/\delta & -\lambda/\delta \\ \lambda^{\star}/\delta & \beta/\delta \end{pmatrix},$$

where $\delta = \sqrt{\beta \beta^* + \lambda \lambda^*}$. The result is the matrix

$$egin{pmatrix} \delta & 0 \ lphaeta^\star/\delta & -lpha\lambda/\delta \end{pmatrix}$$
 .

Since λ and α are not zero (they were diagonal elements of H(k-1)), we know that δ and $-\alpha\lambda/\delta$, the two diagonal elements in the matrix, are also not zero. Note that since the rest of Q is the identity matrix none of the other diagonal elements is affected by the multiplication. Thus, at the end of Step 2, all diagonal elements are non-zero. Since Hermite reduction doesn't introduce any new zeros on the diagonal, the end result of the k-th iteration has all non-zero diagonal elements. But this contradicts the hypothesis on k and our assumption that r < n - 1 was false. Note that for r = n - 1in order to have $h_{n-1,n-1}(k) = 0$, we must have $h_{n,n-1}(k-1) = 0$ and $h_{n-1,n-1}(k-1) \neq 0$.

Next we show that a relation for x must appear as a column of the matrix B. By Lemma 1, $xH_x = 0$. $BA = I_n$ implies $0 = xBAH_x = xBAH_xQ = xBH(k-1)$, where Q is an appropriate unitary $n-1 \times n-1$ matrix. Let z = xB. The above gives

$$(0,\ldots,0) = xBH(k-1) = zH(k-1) = (\ldots,z_{n-1}h_{n-1,n-1}(k-1)).$$

Since $h_{n-1,n-1}(k-1) \neq 0$ then $z_{n-1} = 0$. Hence the n-1-th column of B is a relation for x. \Box

Lemma 6. At any k-th iteration of the algorithm the diagonal entries of H(k) satisfy the inequality $|h_{i,i}(k)| \leq 1$.

Proof. We follow the α, β, λ definitions of the proof of Lemma 5 and use induction. For k = 1 the diagonal entries of H(k) are those of H_x and $s_{j+1} \leq s_j \leq 1$ gives the required inequality. Assume that the inequality also holds up to k-1. The diagonal entries of H(k) are equal to those of H(k-1)except for row r where Step 1 Exchange occurs. When r = n - 1, after the exchange, the r-th diagonal element is β . But $|\beta| \leq |\alpha|/\rho \leq 1$ because $\rho > 1$ and $|\alpha| \leq 1$ by induction. When r < n - 1, after the exchange the r-th diagonal element is δ . But $|\delta| = |\alpha|t \leq 1$ since t < 1 and $|\alpha| \leq 1$. The r + 1-th diagonal element of H is $-\alpha\lambda/\delta$ (as in the proof of Lemma 5) so that $|-\alpha\lambda/\delta| = |\lambda|/t \leq |\alpha|$ because $|\lambda|^2 < |\lambda|^2 + |\beta|^2$ and $|\lambda| \leq |\alpha|t$. \Box

We show that every iteration of PSLQ causes a geometric monotonic increase in a certain function $\Pi(k)$ which is roughly the product of all the principal minors of the matrix H(k). If a relation for x exists, this product will be bounded above and below. Assume x has some relation and as usual let M_x denote the norm of a smallest relation for x. We will need the following technical lemma in the proof of Lemma 9.

Lemma 7. Consider the quotient

$$q(A, B, t) = \frac{\min\{B, t\} \cdot \min\{A, 1\}}{\min\{B, 1\} \cdot \min\{A, t\}}$$

Suppose that the four positive real numbers A, B, 1, t satisfy the three inequalities

 $A\geq B,\quad A\geq t,\quad 1\geq t.$

Then,

$$q(A, B, t) \ge 1.$$

Proof. Of the 16 possible choices in the min's, the inequality $A \ge t$ removes 8, $A \ge B$ removes 2, and $1 \ge t$ removes 1 leaving 5. These five are $A \ge B \ge 1 \ge t$ with quotient $t/1 \cdot 1/t = 1$, $A \ge 1 \ge B \ge t$ with quotient $t/B \cdot 1/t \ge t/1 \cdot 1/t = 1$, $1 \ge A \ge B \ge t$ with quotient $t/B \cdot A/t = A/B \ge 1$, $1 \ge A \ge t \ge B$ with quotient $B/B \cdot A/t = A/t \ge 1$, $A \ge 1 \ge t \ge B$ with quotient $B/B \cdot 1/t = 1/t \ge 1$. \Box

Lemma 8. For α , γ , M_x as above,

$$\gamma^{n-2}M_x|\alpha| \ge 1.$$

Proof. By the choice of r in Step 1 Exchange, we have $\gamma^r |\alpha| \ge \gamma^j |h_{j,j}|$ for any $j, \quad 1 \le j \le n-1$, which implies

$$\gamma^{n-1}/|h_{j,j}| \ge \gamma^r/|h_{j,j}| \ge \gamma^j/|\alpha| \ge \gamma^1/|\alpha|,$$

for all j including that j_o for which $M_x \ge 1/|h_{j_o,j_o}|$ from Theorem 1. Thus $\gamma^{n-2}M_x \ge 1/|\alpha|$ and $\gamma^{n-2}M_x|\alpha| \ge 1$

Definition 6: (The II function). Recall $\tau = \sqrt{1/\rho^2 + 1/\gamma^2}$. Define

$$\Pi(k) = \prod_{1 \le j \le n-1} \min\{\gamma^{n-1} M_x, 1/|h_{j,j}(k)|\}^{n-j}.$$

Lemma 9. For any k > 1 we have (i)

$$(\gamma^{n-1}M_x)^{\binom{n}{2}} \ge \Pi(k) \ge 1,$$

(ii)

$$\Pi(k) \ge \tau \Pi(k-1).$$

Proof. For the k's so far, $h_{j,j}(k) \neq 0$ for all $1 \leq j \leq n-1$. $M_x \geq 1$ and $1/|h_{j,j}(k)| \geq 1$ by Lemma 6. This gives

$$\min\{M_x, 1/|h_{j,j}(k)|\} \ge 1,$$

for all $1 \leq j \leq n-1$, which implies the right hand inequality of (i). On the other hand, it is always the case that $M_x \geq \min\{M_x, 1/|h_{j,j}(k)|\}$, which together with the fact that $\binom{n}{2} = n-1 + \cdots + 2 + 1$ and that $\gamma \geq 1$ gives the left hand inequality of (i).

The proof of part (ii) is more involved. Let r be given by the Step 1 Exchange of PSLQ. Recall the definitions of the two successive diagonal elements α, λ , the single off diagonal element β , $t = \sqrt{\beta\beta^* + \lambda\lambda^*}/|\alpha|$ in the Step 2 (Corner development) of the unitary matrix in terms of β and λ .

Suppose that r < n - 1. Then only two diagonal elements change. These correspond to the 2×2 submatrix of H

$$\begin{pmatrix} \alpha & 0 \\ \beta & \lambda \end{pmatrix}$$

which after a single iteration becomes

$$\begin{pmatrix} \delta & 0\\ \alpha\beta^{\star}/\delta & -\alpha\lambda/\delta \end{pmatrix}.$$

But $|\delta| = |\alpha|t$ so that the absolute values of the of the α, λ diagonal elements are replaced by the absolute values of the $\delta, -\alpha\lambda/\delta$ diagonal elements. All the factors of $\Pi(k)$ are the same except these two so that

$$\frac{\Pi(k)}{\Pi(k-1)} = \left(\frac{\min\{\gamma^{n-1}M_x, 1/(|\alpha|t)\}}{\min\{\gamma^{n-1}M_x, 1/|\alpha|\}}\right)^{n-r} \cdot \left(\frac{\min\{\gamma^{n-1}M_x, t/|\lambda|\}}{\min\{\gamma^{n-1}M_x, 1/|\lambda|\}}\right)^{n-r-1}.$$

Set

$$A = \gamma^{n-1} M_x |\alpha| t \quad and \quad B = \gamma^{n-1} M_x |\lambda|,$$

so that

$$\frac{\Pi(k)}{\Pi(k-1)} = \left(\frac{\min\{A,1\}}{\min\{A,t\}}\right) \cdot \left(\frac{\min\{B,t\}}{\min\{B,1\}} \cdot \frac{\min\{A,1\}}{\min\{A,t\}}\right)^{n-r-1}$$

We now show that the assumptions for Lemma 7 hold. Note that 1 > t by the definition of t; also, $A \ge B$ since $|\alpha|t \ge |\lambda|$. By Lemma 8 we have $A \ge t\gamma \ge t$. By Lemma 7 we have

$$\frac{\Pi(k)}{\Pi(k-1)} \ge \frac{\min\{A,1\}}{\min\{A,t\}} \ge \frac{1}{t} \ge \tau.$$

Now suppose that r = n - 1. By Step 3 Reduction, under one iteration the absolute value of the last diagonal element α is less than $|\alpha|\rho$. All the factors of $\Pi(k)$ except the last are the same so that

$$\frac{\Pi(k)}{\Pi(k-1)} \le \frac{\min\{\gamma^{n-1}M_x, 1/(|\alpha|\rho)\}}{\min\{\gamma^{n-1}M_x, 1/|\alpha|\}} = \frac{\min\{A, t/\rho\}}{\min\{A, t\}}.$$

But we always have $\gamma^{n-2}M_x|\alpha| \ge 1$, so if $A \ge t/\rho \ge t$

$$\frac{\Pi(k)}{\Pi(k-1)} \ge 1/\rho \ge \tau.$$

By Lemma 8, $A \ge t\gamma \ge t$. If $t \le A \le t/\rho$ then

$$\frac{\Pi(k)}{\Pi(k-1)} \ge A/t \ge \gamma \ge \tau.$$

Thus for $r \leq n-1$, $\Pi(k) \geq \tau \Pi(k-1)$. \Box

Theorem 2. Assume real or complex numbers, $n \ge 2$, $\tau > 1$, and that $0 \ne x \in \mathbb{K}^n$ has $\mathbb{O}(\mathbb{K})$ integer relations. Let M_x be the least norm of relations for x. Then $PSLQ(\tau)$ will find some integer relation for x in no more than

$$\binom{n}{2} \frac{\log\left(\gamma^{n-1}M_x\right)}{\log\tau}$$

iterations.

Proof. Suppose we have done k iterations, then from Lemma 6 and Lemma 7, $|h_{j,j}(k)| \neq 0$ and not all $|h_{j,j}(l)| < 1/M_x$ for l < k. By Lemma 6, $\Pi(0) \ge 1$ and by Lemma 7, $\Pi(k) \ge \tau^k$ so that

$$(\gamma^{n-1}M_x)^{\binom{n}{2}} \ge \tau^k$$

Taking natural logarithms of both sides of this inequality gives

$$\binom{n}{2}\log\left(\gamma^{n-1}M_x\right) \geq k\log\tau. \qquad \Box$$

Corollary 2. Let \mathbb{K} be the real numbers \mathbb{R} or the complex numbers \mathbb{C} . Fix n > 1 and assume given a unit n-tuple $x \in \mathbb{K}^n$ which has a relation $m_x \in \mathbb{O}(\mathbb{K})^n$ of least norm M_x . Then there exists a γ such that the algorithm $PSLQ(\tau)$ will construct some $\mathbb{O}(\mathbb{K})^n$ relation for x in no more than

$$2 \cdot (\dim_{\mathbb{R}} \mathbb{K}) \cdot (n^3 + n^2 \log M_x)$$

iterations.

Proof. Let $\gamma = 2$. Then for either \mathbb{K} , $\tau > 1$, specifically, $1/\log \tau < 4 \dim_{\mathbb{R}} \mathbb{K}$. PSLQ(τ) takes O(n) exact arithmetic operations per iteration, so in this sense finds relations in 'polynomial time' $O(n^4 + n^3 \log M_x)$. \Box

5. Upper Bounds on Integer Relations

We compare the relation found by PSLQ to a shortest possible relation.

Lemma 10. Suppose *m* is the relation found on the k + 1-st iteration so that $h_{n-1,n-1}(k+1) = h_{n,n-1}(k) = 0$ and $h_{n-1,n-1}(k) \neq 0$. Then

$$|m| = 1/|h_{n-1,n-1}(k)|.$$

Proof. At this iteration we have developed the matrix $A \in GL(n, \mathbb{O}(\mathbb{K}))$ where the (n-1)-st column of A^{-1} by Lemma 5 is m and the vector $Am^* = e_{n-1}$ has as its only non-zero entry a 1 in the (n-1)-st position. Since $AP = TDQ, Qm^* = D^{-1}T^{\dagger}Am^*$, where T^{\dagger} is the generalized inverse of Tand D is a diagonal matrix with last entry $h_{n-1,n-1}(k)$, which is also the last entry of $D^{-1}T^{\dagger}Am^*$. Because Q is unitary $|Qm^*| = |m^*|$. \Box

Theorem 3. Let M_x be the smallest possible norm of any relation for x. Let m be any relation found by $PSLQ(\tau)$. For all $\gamma > \sqrt{4/3}$ for real vectors and for all $\gamma > \sqrt{2}$ for complex vectors

$$|m| \le \gamma^{n-2} M_x.$$

Proof. Assume we are at the k-th step of PSLQ where a Step 1 Exchange r = n - 1 was made with $h_{n-1,n-1}(k) \neq 0$ and $h_{n-1,n-1}(k+1) = 0$. Then

$$|\gamma^{n-1}|h_{n-1,n-1}(k)| \ge \gamma^j |h_{j,j}(k)|$$

for all $1 \leq j \leq n-2$ by the choice of r. Hence, by Theorem 1, Lemma 8 and Lemma 10

$$M_x \ge 1/\max|h_{i,i}(k)| \ge \gamma^{2-n}/|h_{n-1,n-1}(k)| = \gamma^{2-n}|m|.$$

Comment on Theorem 3. For n = 2, Theorem 3 proves that any relation $0 \neq m \in \mathbb{O}(\mathbb{K}^2)$ found has norm $|m| = M_x$. In other words, $PSLQ(\tau)$ finds a shortest relation. For real numbers this corresponds to the case of the Euclidean algorithm, [13, Book X], [20], [26]. For complex numbers this corresponds to the case of an algorithm in [33].

For n = 3, let x = (113, 343, 311). This vector has a shortest relation $m_x = (7, -15, 14)$ with the shortest norm $|m_x| = M_x = 21.6794...$ This can be verified directly, cf., [25], [31], [11]. On the other hand, for $\tau = 1.0000..., \gamma = 1.1547..., \text{PSLQ}(\tau)$ in iteration 6 produces the relation $m_1 = (24, -7, -1)$. Indeed

$$M_x < |m_1| = 25.0199 \dots \le \gamma M_x = 25.0333 \dots$$

This relation appears from a zero in the second coordinate of the xA_6^{-1} vector. Continuing to iteration 8 gives the relations appearing from the first and second coordinates of the current xA_8^{-1} vector, $m_2 = (-17, -8, 15)$ and $m_3 = (41, 1, -16)$ of norms 24.0416... and 44.0227..., respectively. The vector m_2 has smaller but not smallest norm. Continuing to iterations 9 and 10 gives the relations appearing from the first and second coordinates of xA_9^{-1} of $m_4 = (7, -15, 14)$ and $m_2 = (-17, -8, 15)$, so a shortest vector m_4 was eventually found. In iteration 11 the $h_{2,2}(11) = 0$ condition appeared for the first time giving the relation $m_5 = (-10, -23, 29)$ of norm 38.3405....

This example is instructive in that various choices of the parameter τ give different outputs. The 'legal' τ are such that $1 < \tau < 2$, although the

PSLQ(τ) sometimes works for 'illegal' τ outside of this interval. For the 'legal' τ , $\tau = 1.1$, iteration 6 yields m_1 , 8 yields m_2, m_3 , 9 yields m_4, m_2 , and 10 yields m_5 . On the other hand, for $\tau = 1.8$, iterations 4, 5, 6 all yield only the shortest length relation m_4 . For the 'illegal' τ below 0.7 and above 2.1 the algorithm cycles indefinitely. The end point $\tau = 1.0$ gives essentially the same outputs as $\tau = 1.1$. The other end point $\tau = 2.0$ yields two new relations $m_6 = (1, -91, 100)$ and $m_7 = (0, -311, 343)$ of norms 135.2109... and 463.0010..., respectively.

6. Multiple Relations.

A given unit vector $x \in \mathbb{K}^n$ may have 0, 1, 2, or up to n-1 relations. Once a relation has been constructed, one of the coordinates of xB for the appropriate $B \in GL(n, \mathbb{O}(\mathbb{K}))$ will be zero, and the corresponding column of B will be a relation. The remaining n-1 coordinates can be used to form a new unit vector in $y \in \mathbb{K}^n$. Apply $PSLQ(\tau)$ to this y. Any second relation so found will be integrally independent from the first and can be referred back to the original x. In this way as many as n-1 integrally independent relations for x can be constructed. We omit here the tangent discussion of using classical lattice reduction techniques to find integer relations; this is the case for the Recognize[] function in MathematicaTM which calls the function LatticeReduce[], cf., [11], [12], [27]. Lattice reduction there applies typically only to integer relations for *integer* vectors. Integer relations for *real* or *complex number* vectors.

7. Variations of $PSLQ(\tau)$.

The algorithm $PSLQ(\tau)$ as stated may be performed for various 'illegal' τ or 'illegal' γ , and under these circumstances will find relations for some x vectors. This can happen for $\gamma < \sqrt{4/3}$ in the real case, for $\gamma < \sqrt{2}$ in the complex case, and for $\gamma < \infty$ in the quaternion case, so that $\tau < 1$ and the conclusions of Theorem 2 or Theorem 3 make no sense or have no apparent content. The reason for this apparent anomaly is that for a specific *n*-tuple x the actual field or division ring constant ρ bound in Lemma 4 is not universal and could depend upon an input vector x. Say ρ_x gives a bound such as that of Lemma 4 for some special x or collection of them. Then there may be an "illegal" γ so that $\tau_x = 1/\sqrt{1/\rho^2 + 1/\gamma^2} > 1$. For such x one could expect to see some relation emerge before the number of iterations indicated by Theorem 2 for this $\tau_x = \tau$.

On the other hand, it is possible to use the real $PSLQ(\tau)$ algorithm to find

complex and quaternion relations at the expense of doubling and quadrupling the dimension. For example, suppose z = x + yi + uj + vk is a vector in \mathbb{H}^n with vector components $x, y, u, v \in \mathbb{R}^n$. Suppose the corresponding relation is m = a + bi + cj + dk which is a lattice point in \mathbb{W}^n with integral vector components $a, b, c, d \in \mathbb{Z}^n$. Then $zm^* = 0$ implies four integer relations among the interlaced and suitably sign changed coordinates of z. For the first set $\sum_{1 \leq j \leq n} (a_j x_j - b_j y_j - c_j u_j - d_j v_j) = 0$ and one can apply real PSLQ(τ) to the real 4n-tuple $(\ldots, x_j, y_j, u_j, v_j, \ldots)$. There are three others which are similar. A relation for z will be in the intersection of the four associated lattices. Alternatively, one can give a PSLQ(τ) algorithm along the lines of [23, Section 5. Finding simultaneous integer relations].

8. Computer Implementation of $PSLQ(\tau)$

The PSLQ(τ) algorithm can be implemented using ordinary floating point arithmetic on a computer. Using double precision (i.e., 64-bit) arithmetic, relations of two or three digits in size can be recovered for n up to five or so. Beyond this level, precision is quickly exhausted, and recovered relations and norm bounds are meaningless. Thus a serious implementation of PSLQ (or any other integer relation algorithm for real numbers) must employ some form of multiprecision arithmetic. The authors employed the MPFUN multiprecision translator and computation package. The Fortran-77 version of this software is described in [6], and the newer Fortran-90 version is described in [7]. A C++ translator that employs these routines is also now available. Alternatively, one may employ the multiprecision facilities of symbolic math software packages, such as Maple, Pari or MathematicaTM.

The descriptions presented here of computer implementation of $PSLQ(\tau)$ are for the case of the real number system. Extensions to the case of the complex and quaternions number systems are straightforward, provided one's multiprecision system supports these datatypes.

One key to an efficient implementation is to utilize a simplified version of Hermite reduction and the associated update. As noted in Lemma 3 above, Hermite reduction can be done more efficiently by a triply nested loop. In fact, the update operations associated with Hermite reduction (updating x, H, A and B) can also be done in a loop of this form. Further, if these updates are done in this manner, then it is not necessary to compute the Dmatrix. This simplified scheme is as follows. In the initialization step, Hermite reduction and the subsequent updates are replaced with the following:

For *i* from 2 to *n*, for *j* from i - 1 to 1 (step -1), set $t = \text{nint}(h_{i,j}/h_{j,j})$ and replace x_j by $x_j - tx_i$; then for *k* from 1 to *j* replace $h_{i,k}$ by $h_{i,k} - th_{j,k}$; for

k from 1 to n replace $a_{i,k}$ by $a_{i,k} - ta_{j,k}$ and replace $b_{k,j}$ by b(k,j) + tb(k,i).

Step 3 is also replaced with this, except i is incremented from r+1 to n, and j is decremented from min $\{i-1, r+1\}$ to 1. Here r denotes the row index selected in Step 1. These more restrictive limits on i and j merely reflect the fact that t = 0 outside these limits.

Obviously in a computer implementation some care must be taken in testing for zero. This is typically done by checking that the absolute value of the tested value is less than the "epsilon" appropriate for the level of numeric precision being used. Also, a run should be terminated if any entry of the *A* matrix exceeds the level of numeric precision being used (so that these integer values can no longer be represented exactly).

The level of working precision required for PSLQ is generally only a few digits greater than the accuracy of the input x vector. Along this line, if one wants to recover (or to exclude) relations of size d digits, then the input data must be specified to at least nd digits in order to obtain numerically meaningful results. The significance of a recovered result can be measured by noting the ratio between the multiprecision epsilon and the largest entry of the updated x vector when a relation is recovered. If this ratio is very small, such as 10^{-40} , then one can be fairly certain that the relation produced by PSLQ is a real relation. But if this ratio is only a few orders of magnitude below unity, then the result is suspect, and higher accuracy in the input data, as well as correspondingly higher working precision, is required.

The above implementation is satisfactory for most applications. For more demanding applications, a "two-level" implementation is significantly faster. In a two-level implementation, most operations are performed in ordinary double precision arithmetic, with occasional updates of multiprecision arrays using multiprecision arithmetic. This two-level scheme can be described as follows. Here the prime notation is used to denote double precision approximations to multiple precision values.

To initialize, perform the initialization step as described above using full precision. Then perform a "double precision initialization": (1) set $x' = x/\max_{i,j} |x_j|$ and set H' = H; (2) perform a LQ decomposition on H', using double precision arithmetic, setting H' to be the lower triangular part; (3) set $A' = B' = I_n$.

PSLQ iterations are then performed as above on the arrays x', H', A'and B', using double precision arithmetic. Some care must be taken to insure numerical accuracy in these iterations. Obviously these iterations before entries in A' grow so large (9 × 10¹⁵ on IEEE systems) that they cannot be exactly represented as double precision values. In the authors'

implementation, double precision iterations are halted when the largest entry of A' exceeds 10^{10} . Tests for zero in these iterations must reflect the accuracy of double precision arithmetic — the authors used an "epsilon" of 10^{-13} here. As an additional measure to insure numerical integrity, the authors' code aborts the modified Hermite reduction procedure (and restores arrays to their previous values) if the multiplier q exceeds 10^7 .

When the double precision iterations are halted, either due to large entries in A', or to a tentative zero in x' or H', it is necessary to perform a "multiprecision update": (1) replace A by A'A, replace B by BB', replace H by A'H, and replace x by xB'; (2) check for zero entries in x, using the multiprecision epsilon. If no zeroes are found, then a double precision initialization is performed, followed by more double precision PSLQ iterations.

One detail has been omitted here. In some cases, the entries of the updated x vector have such a large dynamic range (greater than 10^{10} in the authors' implementation) that when converted to double precision, additions and subtractions would produce results of questionable reliability. In these cases it is necessary to perform PSLQ iterations on the multiprecision arrays, using multiprecision arithmetic, for a number of iterations until this large dynamic range is eliminated. If this situation is encountered on any iteration other than the very first, a multiprecision LQ decomposition of Hmust be performed prior to performing these multiprecision iterations (so that the H array contains the same entries as the H array defined in the PSLQ algorithm statement).

The authors' Fortran implementation of PSLQ, together with the required multiprecision arithmetic software, is available by sending electronic mail to dbailey@nas.nasa.gov. Also available are MathematicaTM implementations of PSLQ as well as a number of other integer relation algorithms for comparison.

9. Summary of the Literature

The problem of finding integer relations among sets of rational and real numbers is quite old. When n = 2 this problem can be solved for rationals by the first Euclidean algorithm in Euclid, Book VII, and for reals by the second Euclidean algorithm given in Euclid, Book X, cf., [26], [11], [37]. Generalizations of this algorithm to higher real dimensions were proposed without proof by many authors, including Jacobi [24], Hermite [22], Poincaré [32], Perron [30], Brun [9, 10] and Szekeres [38]. Various counterexamples can be found in [15] and [19].

The first integer relation finding algorithm with proofs for the case of real numbers was discovered in 1977 by Ferguson and Forcade, [14, 15]. These

algorithms were shown to be polynomial time in the logarithm of the size of a smallest relation. They were not shown to be polynomial in the dimension. Since then, other related algorithms for finding relations for real vectors have appeared in [8], [16], [17], [18]. For example, [5] reports on a computer implementation of [16]. The sequence including [23] (HJLS), [2] and [1] (PSLQ), [3] (a concise statement of PSLQ), and [35] (a stable variation of HJLS) will be discussed below.

These algorithms all depend upon an orthogonal decomposition of some kind. See [21], for a list of various orthogonalization algorithms and their numerical linear algebra differences. PSLQ is of the QR type. HJLS follows the lattice reduction work of [28], [34], and [36], which is classical Gram-Schmidt type, cf. [31] and [11]. This conceptual difference may explain some of the numerical differences observed between PSLQ and HJLS, cf. [2].

Rigorous proofs that the algorithm under investigation must find a relation if one exists appeared in [14, 8, 15, 16]. All of these proofs gave a linear bound in the logarithm of the size of a relation, but were not known to be polynomial in the dimension. [8] and [16] had unsatisfactory proofs in the sense that they were shown to be at worst exponential in the dimension rather than polynomial in the dimension. This unsatisfactory state of affairs was resolved affirmatively with the proofs that appeared in [23] for the 'small integer relation algorithm'. We will refer to this 'small integer relation algorithm' as HJLS, as stated in [23, Section 3] as a reflection of that in [8, Section 3]. In fact, this proof in [23] was the first appearance in the literature of a 'polynomial time' bound for a relation finding algorithm, polynomial in both dimension and logarithm of relation size.

This important progress was made when [23] combined two independent streams of research, [14, 8, 15, 16, 18] and [28, 29, 34, 35, 11]. Inspired by the polynomial result of [23], but not the details, the first author of this paper formulated what he thought was a new algorithm [2, 1] and gave a polynomial proof. This proof was independent of that of [23], a different analysis, but flawed by giving a slightly higher degree polynomial in the dimension than the polynomial proof given in [23]. This algorithm in [2, 1] was called PSLQ and had the advantage of the adjustable parameter γ or τ . Applications and implementation of this earlier version of PSLQ(τ) were described in [3, 7, 4]. These implementations showed that the parameters were a helpful feature of the algorithm. The bound on iterations for HJLS proven in [23] was $O(n^3 + n^2 \log_2 M_x)$; this is consonant with the bound proven in this paper for PSLQ($\sqrt{2}$). The subsequent paper [35] included parameters as well as addressing a certain issue of stability.

As a specific example, consider the triple x = (11, 27, 31). We list the

sequence of A^{-1} matrices for each algorithm. A relation if found will be constructed as a column of one of these A^{-1} matrices.

For PSLQ(1.1547) the successive iterations k = 0, 1, 2, 3, 4, yield the five A^{-1} matrices

$$\begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & -1 & 1 \end{pmatrix}, \begin{pmatrix} 1 & 0 & 0 \\ 3 & 8 & 1 \\ -3 & -7 & -1 \end{pmatrix}, \begin{pmatrix} -2 & 1 & 0 \\ 2 & 3 & 1 \\ -1 & -3 & -1 \end{pmatrix}, \begin{pmatrix} 3 & -2 & 0 \\ 1 & 2 & 1 \\ -2 & -1 & -1 \end{pmatrix}, \begin{pmatrix} -1 & -8 & -2 \\ 5 & 9 & 2 \\ -4 & -5 & -1 \end{pmatrix}.$$

Note that PSLQ has constructed two relations appearing as the first and second columns of the last matrix, iteration k = 4.

For HJLS the successive iterations k = 0, 1, 2, 3, 4, 5, 6 yield the seven A^{-1} matrices

$$\begin{pmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix}, \begin{pmatrix} 1 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 0 & 1 & 0 \\ 0 & 0 & 1 \\ 1 & 0 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}, \begin{pmatrix} 1 & -2 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & -1 \end{pmatrix}.$$

Note that only one relation is found; it appears in the last column of the last matrix, iteration k = 6. The authors of [34] claimed that HJLS is a special case of $PSLQ(\tau)$ for $\gamma = \sqrt{2}$ or equivalently $\tau = \sqrt{4/3}$. The example just given shows that this claim cannot be true.

The significance of the parameter was revealed clearly in the extensive tables appearing in [2]. In [2] the parameter was γ which is equivalent to giving $\tau = 1/\sqrt{\rho^2 + 1/\gamma^2}$. The choice of the parameter τ has precision consequences: depending upon the choice of parameter a numerical precision much higher than that of the input real vector must be used to obtain a reliable result. For example, the algebraic number

$$\alpha = 3^{1/4} - 2^{1/4} = \sqrt{\sqrt{3}} - \sqrt{\sqrt{2}}$$

satisfies a polynomial of degree 16 with coefficients

$$(1, 0, 0, 0, -3860, 0, 0, 0, -666, 0, 0, 0, -20, 0, 0, 0, 1).$$

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The algorithm $PSLQ(\tau)$ for $\tau = 1.000006145$ or $\gamma = 1.1547005384$, applied to the vector $(1, \alpha, \alpha^2, \dots, \alpha^{n-1})$, with n = 17, finds these coefficients with a working precision of 75 decimal digits. We have shown with the n = 3example above that HJLS is not $PSLQ(\sqrt{4/3})$. Again, we see that HJLS requires a working precision of more than 10,000 decimal digits to find this n = 17 relation. Comparative run times are not particularly relevant here but are also correspondingly higher for HJLS — see Table 2 of [2].

For a slightly different $\tau = \sqrt{4/3} = 1.154700538...$, PSLQ(τ) requires 85 decimal digits, 10 digits more than for $\gamma = 1.1547005384$. Generally the closer τ is to 1 the less precision seems to be required. This observed phenomenon appears to have nothing to do with any question of numerical stability.

The various algorithms in the literature stand independently of their published proofs; their published proofs may not reveal their actual properties clearly. Though the proofs were exponential, the algorithms stated in [14], and in [15], and again in [16] were parametric. The parameter b in [14, 15] satisfies 1 < b < 2 whereas in [16] the parameter γ is emphasized. The algorithm in [8, Sect. x] seems closest to $PSLQ(\sqrt{4/3})$ with the τ parameter set by $\gamma = \sqrt{2}$. This parameter choice appears in [8, Sect. x] without the [28] setting and reappears in [23] as the "small integer relation algorithm", which we call HJLS, rewritten in the [28] language and accompanied by a 'polynomial time' proof for the first time.

Bergman discussed the complex case of finding gaussian integer relations for complex vectors in [8, Sect. 5: Variants]. Bergman also gave an algorithm for the simultaneous real vector case in [8, Sect. 7]. Following Bergman, the paper defining HJLS for simultaneous real vectors, [23, cf., Sect. 5], implicitly includes the complex and quaternion vector case as well. As an alternate approach, inspired by [37], in this paper we have extended the base field of $PSLQ(\tau)$ to these division rings and introduced unitary matrices into the algorithm directly. The proof given here of polynomial number of iterations covers the real and complex cases, but fails for quaternions. However, the quaternion version of $PSLQ(\tau)$ performs reasonably well experimentally in finding hamiltonian integer relations for quaternion vectors. This was explained in Section 8.

10. Open Questions

1) Is there a relation finding algorithm that finds a shortest relation in a polynomial (in the dimension) number of iterations?

2) What are the best choices for the parameter τ or γ relative to the

number of iterations, time, and precision requirements of PSLQ?

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