HOW TO PREDICT CONGRUENTIAL GENERATORS

by

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ABSTRACT

In this paper we show how to predict a large class of pseudorandom number generators. Namely, generators outputting a sequence of integers \( s_0, s_1, \ldots \) where \( s_i \) is computed by the recurrence

\[
s_i = \sum_{j=1}^{k} \alpha_j \Phi_j(s_0, s_1, \ldots, s_{i-1}) \pmod{m}
\]

for integers \( m \) and \( \alpha_j, \) and integer functions \( \Phi_j, j = 1, \ldots, k. \) Our predictors are efficient, provided that the functions \( \Phi_j \) are computable (over the integers) in polynomial time. These predictors have access to the elements of the sequence prior to the element being predicted, but they do not know the modulus \( m \), nor the coefficients \( \alpha_j \) the generator actually works with. This extends previous results about the predictability of such generators. In particular, we prove the predictability of multivariate polynomial generators, i.e. generators where \( s_i \equiv P(s_{i-n}, \ldots, s_{i-1}) \pmod{m}, \) for a polynomial \( P \) of fixed degree in \( n \) variables.

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1. INTRODUCTION

A number generator is a deterministic algorithm that given a sequence of initial values, outputs an (infinite) sequence of numbers. Some generators, called pseudorandom number generators, are intended to output sequences of numbers having some properties encountered in truly random sequences. Such generators appear in diverse applications as Probabilistic Algorithms, Monte Carlo Simulations, Cryptography, etc. For cryptographic applications a crucial property for the sequences generated is their unpredictability. That is, the next element generated should not be predictable, even given the entire past sequence.

A pseudorandom number generator that has received much attention is the so-called linear congruential generator, an algorithm that on input integers \( a, b, m, s_0 \) outputs a sequence \( s_1, s_2, \ldots \) where

\[
s_i = a \cdot s_{i-1} + b \pmod{m}
\]

Knuth [12] extensively studied some statistical properties of these generators.

Boyar [15] proved that linear congruential generators are predictable (in a sense to be formalized later), even when the coefficients and the modulus are unknown to the predictor. Later, Boyar [3] extended her method, proving the predictability of a large family of generators. She considered general congruential generators where the element \( s_i \) is computed as

\[
s_i = \sum_{j=1}^{k} \alpha_j \Phi_j(s_0, s_1, \ldots, s_{i-1}) \pmod{m}
\]  

for some integers \( m \) and \( \alpha_j \), and computable integer functions \( \Phi_j, j=1,\ldots,k \). She showed that these sequences can be predicted, for some class of functions \( \Phi_j \), by a predictor knowing these functions and able to compute them, but not given the coefficients \( \alpha_j \), nor the modulus \( m \). Boyar's method requires that the functions \( \Phi_j \) have the unique extrapolation property. The functions \( \Phi_1, \Phi_2, \ldots, \Phi_k \) have the unique extrapolation property with length \( r \), if for every pair of generators working with the above set of functions, the same modulus \( m \) and the same initial values, if both generators coincide in the first \( r \) values generated by them, then they output the same infinite sequence. Note that these generators need not be identical (i.e., they may have different coefficients).

Boyar's method allows the construction of efficient predictors for generators having the unique extrapolation property with length \( r \), for a small (and known) integer \( r \). The linear congruential generator is an example of a generator having the extrapolation property (with length 2). Boyar proved this property also for two extensions of the linear congruential generator. Namely, the generators in which the element \( s_i \) satisfies the recurrence

\[
s_i = \alpha_1 s_{i-k} + \cdots + \alpha_k s_{i-1} \pmod{m}
\]

and those for which
The first case with length \( k + 1 \), the second with length 3. She also conjectured the predictability of generators having polynomial recurrence:

\[ s_i = p(s_{i-1}) \pmod{m} \]

for an unknown polynomial \( p \) of fixed (and known) degree.

A natural generalization of the above examples is a generator having multivariate polynomial recurrence, that is a generator outputting a sequence \( s_0, s_1, \ldots \) where

\[ s_i = P(s_{i-n}, \ldots, s_{i-1}) \pmod{m} \]

for a polynomial \( P \) in \( n \) variables. Note that for polynomials \( P \) of fixed degree and fixed \( n \), the recurrence is a special case of the general congruential generators. Lagarias and Reeds [14] showed that multivariate polynomial recurrences have the unique extrapolation property. Furthermore, for the case of a one-variable polynomial, they proved this property with length equal to the degree of the polynomial, thus settling Boyar's conjecture concerning the predictability of such generators. However, for the general case they did not give a bound on the length for which these recurrences are extrapolatable (neither a way to compute this length). Thus, unfortunately, Boyar's method does not seem to apply for general multivariate polynomial recurrences.

In this paper we show how to predict any general congruential generator, i.e. any generator of the form (1). The only restriction on the functions \( \Phi_j \) is that they are computable in polynomial time when working over the integers. This condition is necessary to guarantee the efficiency of our method. (The same is required in Boyar's method.) Thus, we remove the necessity of the unique extrapolation property, and extend the predictability results to a very large class of generators. In particular, we show that multivariate polynomial recurrence generators are predictable.

Our predicting technique is based on ideas from Boyar's method, but our approach to the prediction problem is somewhat different. Boyar's method tries to simulate the generator by "discovering" its secrets: the modulus \( m \) and the coefficients \( \alpha_j \) that the generator works with. Instead, our algorithm uses only the knowledge that these coefficients exist, but does not try to find them. Some algebraic techniques introduced by Boyar when computing over the integers, are extended by us to work also when computing over the ring of integers modulo \( m \).
2. DEFINITIONS AND NOTATION

Definition: A number generator is an algorithm that given \( n_0 \) initial values, denoted \( s_{-n_0}, \ldots, s_{-1} \), outputs an infinite sequence \( s_0, s_1, \ldots \) where each element \( s_i \) is computed deterministically from the previous elements, including the initial values.

For example, a generator of the form \( s_i = \alpha_1 s_{i-k} + \ldots + \alpha_k s_{i-1} \pmod{m} \) requires a set of \( k \) initial values to begin computing the first elements \( s_0, s_1, \ldots \) of the sequence. Thus, for this example \( n_0 = k \).

Definition: A (general) congruential generator is a number generator for which the \( i \)-th element of the sequence is computed by the congruence

\[
s_i = \sum_{j=1}^{k} \alpha_j \Phi_j(s_{-n_0}, \ldots, s_{-1}, s_0, \ldots, s_{i-1}) \pmod{m}
\]

where \( \alpha_j \) and \( m \) are arbitrary integers and \( \Phi_j, 1 \leq j \leq k \), is a computable integer function. For a given set of \( k \) functions \( \Phi = \{ \Phi_1, \Phi_2, \ldots, \Phi_k \} \) a congruential generator working with these functions (and arbitrary coefficients and modulus) will be called a \( \Phi \)-generator.

Note that the functions \( \Phi_j \) work on sequences of elements, so the number of arguments for these functions is variable. Some matrix notation will be more convenient.

Notation: \( s(i) \) will denote the vector of elements (including the initial values) until the element \( s_i \), i.e.,

\[
s(i) = (s_{-n_0}, \ldots, s_{-1}, s_0, \ldots, s_{i}) \quad i=-1,0,1,2,\ldots
\]

Thus, \( \Phi_j(s_{-n_0}, s_{-1}, s_0, \ldots, s_{i-1}) \) will be written as \( \Phi_j(s(i-1)) \).

Let \( \alpha \) denote the vector \((\alpha_1, \alpha_2, \ldots, \alpha_k)\) and \( B_i, i \geq 0 \), denote the column vector

\[
B_i = \begin{bmatrix}
\Phi_1(s(i-1)) \\
\Phi_2(s(i-1)) \\
\vdots \\
\Phi_k(s(i-1))
\end{bmatrix}
\]

Then we can rewrite the \( \Phi \)-generator's recurrence as

\[
s_i = \alpha \cdot B_i \pmod{m} \tag{2}
\]

Here, and in the sequel, \( \cdot \) denotes matrix multiplication.

Finally, \( B(i) \) will denote the matrix

\[
B(i) = \begin{bmatrix}
B_0 & B_1 & \cdots & B_i
\end{bmatrix}
\]

For complexity considerations we refer to the size of the prediction problem as given by the size of the modulus \( m \) and the number \( k \) of coefficients the generator actually works with. (Note that the coefficients as well as the elements output by the generator have size at most \( \log m \)). We
consider as efficient generators for which the functions $\Phi_j, 1 \leq j \leq k$, are computable in time polynomial in $\log m$ and $k$. Also the efficiency of a predictor will be measured in terms of these parameters, which can be seen as measuring the amount of information hidden from the predictor.

We shall be concerned with the complexity of the functions $\Phi_j$ when acting on the vectors $s(i)$, but computed over the integers (and not reduced modulo $m$). This will be referred to as the non-reduced complexity of the functions $\Phi_j$. The performance of our predicting algorithm will depend on this complexity.

Definition: $\Phi$-generators having non-reduced time-complexity polynomial in $\log m$ and $k$ are called non-reduced polynomial-time $\Phi$-generators.

Next we define the basic concept, throughout this paper, of a predictor:

Definition: A predictor for a $\Phi$-generator is an algorithm that interacts with the $\Phi$-generator in the following way. The predictor gets as input the initial values that the generator is working with. For $i = 0, 1, 2, \ldots$ the predictor outputs its prediction for the element $s_i$ and the generator responds with the true value of $s_i$.

An efficient predictor (for a $\Phi$-generator) is a predictor for which there exist polynomials $P$ and $Q$ such that

1) the computation time for every prediction is bounded by $P(k, \log m)$
2) the number of prediction mistakes is bounded by $Q(k, \log m)$

Observe that when computing its prediction for $s_i$ the predictor has seen the entire segment of the sequence before $s_i$, and the initial values. The only secret information kept by the generator is the coefficients and the modulus. If the generator is not given the initial values then our method cannot be applied to arbitrary $\Phi$-generators. However, in typical cases (including the multivariate polynomial recurrence) generators have recurrences depending only on the $n_0$ last elements, for some constant $n_0$. In this case the predictor may consider the first $n_0$ elements generated as initial values, and begin predicting after the generator outputs them.
3. THE PREDICTING ALGORITHM

The predictor tries to infer the element \( s_i \) from knowledge of all the previous elements of the sequence, including the initial values. It does not know the modulus \( m \) the generator is working with. So, the predicting algorithm begins by computing over the integers. After some portion of the sequence is revealed, it computes an integer \( \hat{m} \), a multiple of the true \( m \). From this point its computations will be modulo \( \hat{m} \). Later, after a number of mistakes are made by the predictor, \( \hat{m} \) will be updated, and so on. The values of \( \hat{m} \) are always multiples of \( m \), but also (nontrivial) divisors of the former \( \hat{m} \). Eventually \( \hat{m} \) may reach the true value of \( m \). (For degenerate cases, like a generator producing a constant sequence, it may happen that \( m \) will never be reached but this will not effect the prediction capabilities of the algorithm).

We shall divide the predicting algorithm into two stages. The first stage is when working over the integers. The second one is after the first modulus \( \hat{m} \) was computed. The distinction between these two stages is not essential, but some technical reasons make it convenient. In fact, the algorithm is very similar for both stages. For the first, one can think of it as working with modulus \( \hat{m} = \infty \).

The idea behind the algorithm is to find linear dependencies among the columns of the matrix \( B(i) \) and to use these dependencies for the prediction of the next element \( s_i \). More specifically, we try to find a representation of \( B_i \) as a linear combination (modulo the current \( \hat{m} \) ) of the previous \( B_j \)'s (that are known to the predictor at this time). If such a combination exists, we apply it to the previous elements in the sequence (i.e. previous \( s_j \)'s) to obtain our prediction for \( s_i \). If not correct, we made a mistake but gain information that allows us to refine the modulus \( \hat{m} \). A combination as above will not exist if \( B_i \) is independent of the previous columns. We show that under a suitable definition of independence, the number of possible independent \( B_i \)'s cannot be too large. Therefore a small number of mistakes can be guaranteed, allowing us to prove the efficiency of the predictor.

The number of mistakes made by the predictor, until it is able to refine the current \( \hat{m} \), is bounded by a polynomial in the size of this \( \hat{m} \). Also the total number of distinct moduli \( \hat{m} \) computed during the algorithm is bounded by the size of the first (finite) \( \hat{m} \). Thus, the total number of possible mistakes is polynomial in this size, which in turn is determined by the length of the output of the non-reduced functions \( \Phi_j \). In particular, if the non-reduced complexity of these functions is polynomial in the size of the true \( m \) and \( k \) (i.e. we are predicting a non-reduced polynomial time \( \Phi \)-generator) then the total number of mistakes made by the predictor will be also polynomial in these parameters. The same is true for the computation time of every prediction.
It is interesting at this point to compare the present algorithm to Boyar's [3]. Our first stage is exactly the same as the first stage there. That is, the two algorithms begin by computing a multiple of the modulus \( m \). Once this is accomplished, Boyar's strategy is to find a set of coefficients \( \{ \alpha_j \}_{j=1}^J \), that will be refined during the algorithm, together with the moduli \( m_j \), until no more mistakes will be made. For proving the correctness and efficiency of her predictor, it is required that the generator satisfies the \textit{unique extrapolation property} (mentioned in the Introduction). In our work, we do not try to find the coefficients. Instead, we extend the ideas of the first stage, and apply them also in the second stage. In this way the need for an extrapolation property is avoided, allowing the extensions of the predictability results.

3.1 First Stage

Let us describe how the predictor computes its prediction for \( s_i \). At this point the predictor knows the whole sequence before \( s_i \), i.e. \( s(i-1) \), and so far it has failed to compute a finite multiple of the modulus \( m \), so it is still working over the integers. In fact, the predictor is able at this point to compute all the vectors \( B_0, B_1, \ldots, B_i \), since they depend only on \( s(i-1) \). Moreover, our predictor keeps at this point, a submatrix of \( B(i-1) \), denoted by \( \overline{B(i-1)} \), of linearly independent (over the rationals) columns. (For every \( i \), when predicting the element \( s_i \), the predictor checks if the column \( B_i \) is independent of the previous ones. If this is the case then \( B_i \) is added to \( \overline{B(i-1)} \) to form \( \overline{B(i)} \).) Finally, let us denote by \( \overline{s(i-1)} \) the corresponding subvector of \( s(i-1) \), having the entries indexed with the same indices appearing in \( \overline{B(i-1)} \).

Prediction of \( s_i \) in the first stage:

The predictor begin by computing the (column) vector \( B_i \). Then, it solves, over the rationals, the system of equations

\[
\overline{B(i-1)} \cdot x = B_i
\]

If no solution exists, \( B_i \) is independent of the columns in \( \overline{B(i-1)} \) so it sets

\[
\overline{B(i)} = \left[ \overline{B(i-1)} \quad B_i \right]
\]

and it fails to predict \( s_i \).

If a solution exists, let \( c \) denote the solution (vector) computed by the predictor. The prediction for \( s_i \), denoted \( p \), will be

\[
p = \overline{s(i-1)} \cdot c
\]

The predictor, once having received the true value for \( s_i \), checks whether this prediction is correct or not (observe that the prediction \( p \) as computed above may not even be an integer). If
correct, it has succeeded and goes on predicting $s_{i+1}$. If not, i.e. $p \not\equiv s_i$, the predictor has made a mistake, but now it is able to compute $\tilde{m} \neq \infty$, the first multiple of the modulus $m$, as follows. Let the solution $c$ be

$$c = \begin{bmatrix} c_1/d_1 \\ c_2/d_2 \\ \vdots \\ c_l/d_l \end{bmatrix}$$

where $c_j$ and $d_j$ are relatively prime integers; for $j = 1, 2, \ldots, l$. Now, let $d$ denote the least common multiple of the dominators in these fractions, i.e. $d = \text{lcm}(d_1, \ldots, d_l)$. The value of $\tilde{m}$ is computed as follows

$$\tilde{m} = dp - ds_i$$

Observe that this $\tilde{m}$ is an integer, even if $p$ is not. Moreover this integer is a multiple of the true modulus $m$ the generator is working with (see Lemma 1 below).

Once the first $\tilde{m}$ was computed, the predictor can begin working modulo this $\tilde{m}$. So the first stage of the algorithm is terminated and it goes on into the second one.

The main facts concerning the performance of the predicting algorithm during the first stage are summarized in the next Lemma.

Lemma 1:

a) The number $\tilde{m}$ computed at the end of the first stage is a nonzero multiple of the modulus $m$.

b) The number of mistakes made by the predictor in the first stage is at most $k+1$.

c) For non-reduced polynomial time $\Phi$-generators, the prediction time for each $s_i$ during the first stage is polynomial in $\log m$ and $k$.

d) For non-reduced polynomial time $\Phi$-generators, $\tilde{m}$ can be computed such that its size will be polynomial in $\log m$ and $k$.

Proof:

a) From the definition of the generator we have the congruence $s_j \equiv \alpha B_j \pmod{m}$ for all $j \geq 0$, therefore

$$\overline{s(i-1)} \equiv \alpha B \overline{(i-1)} \pmod{m} \quad (3)$$
Thus,

\[ dp = \bar{d} \bar{s}(i-1) \cdot c \]  
(by definition of \( p \))

\[ = d \alpha \cdot B(i-1) \cdot c \pmod{m} \]  
(by (3))

\[ = d \alpha \cdot B_i \]  
\((c \text{ is a solution } B(i-1) \cdot x = B_i)\)

\[ = d s_i \pmod{m} \]  
(By definition of \( s_i \) (2))

So we have shown that \( dp \equiv ds_i \pmod{m} \). Observe that it cannot be the case that \( dp = ds_i \), because this implies \( p = s_i \), contradicting the incorrectness of the prediction. Thus, we proved that \( \delta t = dp - ds_i \) is indeed a nonzero multiple of \( m \).

b) The possible mistakes in the first stage are when a solution to the system of equations \( B(i-1) \cdot x = B_i \) does not exist, or when such a solution exists but our prediction is incorrect. The last case will happen only once because after that occurs the predictor goes into the second stage. The first case cannot occur "too much". Observe that the matrices \( B(j) \) have \( k \) rows, thus the maximal number of independent columns (over the rationals) is at most \( k \). So the maximal number of mistakes made by the predictor in the first stage is \( k+1 \).

c) The computation time for the prediction of \( s_i \) is essentially given by the time spent computing \( B_i \) and solving the above equations. The functions \( \Phi_j \) are computable in time polynomial in \( \log m \) and \( k \), so the computation of the vector \( B_i \) is also polynomial in \( \log m \) and \( k \). The complexity of solving the equations system, over the rationals, is polynomial in \( k \) and in the size of the entries of \( B(i-1) \) and \( B_i \) (see [8], [17, Ch.3]). These entries are determined by the output of the (non-reduced) functions \( \Phi_j \), and therefore their size is bounded by a polynomial in \( \log m \) and \( k \). Thus, the total complexity of the prediction step is polynomial in \( \log m \) and \( k \), as required.

d) As said in the proof of part c), a solution to the system of equations in the algorithm, can be found in time bounded polynomially in \( \log m \) and \( k \). In particular this guarantees that the size of the solution will be polynomial in \( \log m \) and \( k \). (By size we mean the size of the denominators and numerators in the entries of the solution vector.) Clearly, by the definition of \( \delta t \), the polynomiality of the size of the solution \( c \), implies that the size of \( \delta t \) is itself polynomial in \( \log m \) and \( k \). \( \square \)
3.2 Second Stage

After having computed $m$, the first multiple of $m$, we proceed to predict the next elements of the sequence, but now working modulo $m$. The prediction step is very similar to the one described for the first stage. The differences are those that arise from the fact that the computations are modulo an integer. In particular the equations to be solved will not be over a field (in the first stage it was over the rationals), but rather over the ring of residues modulo $m$. Let us denote the ring of residues modulo $n$ by $Z_n$. In the following definition we extend the concept of linear independence to these rings.

**Definition:** Let $v_1, v_2, \ldots, v_l$ be a sequence of $l$ vectors with $k$ entries from $Z_n$. We say that this sequence is *linearly independent mod* $n$ if $v_1 \neq 0$ and there exist no index $i, 2 \leq i \leq l$, and elements $c_1, c_2, \ldots, c_{i-1} \in Z_n$, such that $v_i \equiv c_1 v_1 + c_2 v_2 + \cdots + c_{i-1} v_{i-1} \pmod{n}$.

Note that the order here is important. Unlike the case in the traditional definition over a field, in the above definition it is not equivalent to say that no vector in the set can be written as a linear combination of the others. Another important difference is that it is not true in general, that $k+1$ vectors of $k$ components over $Z_n$ must contain a dependent vector. Fortunately, a slightly weaker statement does hold.

**Lemma 2:** Let $l, k, n$ be integers, and $v_1, v_2, \ldots, v_l$ be a sequence of $k$-dimensional vectors over $Z_n$. If the sequence is linearly independent mod $n$ then $l \leq k \log_q n$, where $q$ is the smallest prime dividing $n$.

**Proof:** Let $v_1, v_2, \ldots, v_l$ be a sequence of $l$ vectors from $Z_n^k$, and suppose this sequence is linearly independent mod $n$. Consider the set

$$V = \{ \sum_{i=1}^l c_i v_i \pmod{n} : c_i \in \{0, 1, \ldots, q-1\} \}$$

We shall show that this set contains $q^l$ different vectors. Equivalently, we must show that no two combinations in $V$ yield the same vector. Thus we must prove the following

**Claim:** For every $c_i, c'_i \in \{0, 1, \ldots, q-1\}, 1 \leq i \leq l$, if $\sum_{i=1}^l c_i v_i \equiv \sum_{i=1}^l c'_i v_i \pmod{m}$ then $c_i = c'_i$ for $i=1, 2, \ldots, l$.

Suppose this is not true. Then we have $\sum_{i=1}^l (c_i - c'_i) v_i \equiv 0 \pmod{n}$. Denote $c_i - c'_i$ by $d_i$. Let $l$ be the maximal index for which $d_i \neq 0$. This number $d_i$ satisfies $-q < d_i < q$, so it has an inverse modulo $n$ (recall that $q$ is the least prime divisor of $n$), denoted $d_i^{-1}$. It follows that $v_l \equiv -d_i^{-1} d_i v_i \pmod{n}$ contradicting the independence of $v_i$, and thus proving the claim.
Hence, \(|V| = q^l\) and therefore

\[ q^l = |V| \leq |Z_q^k| = n^k \]

which implies \(l \leq k \log_q n\), proving the lemma. \(\Box\)

With the above definition of independence in mind, we can define the matrix \(\overline{B}(i)\) as a submatrix of \(B(i)\), in which the (sequence of) columns are linearly independent \(\mod m\). Note that \(m\) will have distinct values during the algorithm, so when writing \(\overline{B}(i)\) we shall refer to its value modulo the current \(m\).

**Prediction of \(s_i\) in the second stage:**

Let us describe the prediction step for \(s_i\) when working modulo \(m\). In fact, all we need is to point out the differences with the process described for the first stage.

As before, we begin by computing the vector \(B_i\) (now reduced modulo \(m\)), and solving the system of equations

\[ \overline{B}(i-1) \cdot x \equiv B_i \pmod{m} \]

We stress that this time we are looking for a solution over \(Z_m\). In case a solution does not exist, we fail to predict, exactly as in the previous case. As before the vector \(B_i \pmod{m}\) is added to \(\overline{B}(i-1)\) to form the matrix \(\overline{B}(i)\). If a solution does exist we output our prediction, computed like before, but the result is reduced mod \(m\). Namely, we set

\[ p = s(i-1) \cdot c \pmod{m}, \]

where \(c\) is a solution to the above system of modular equations. If the prediction is correct, we proceed to predict the next element \(s_{i+1}\). If not, we take advantage of this error to update \(m\).

The new \(m\) is computed as follows. Let \(c = (c_1, \ldots, c_i) \in Z_m^i\) be the computed solution for the system of equations and \(p\) the prediction for \(s_i\). Set

\[ m' = \gcd(m, p-s_i) \]

This \(m'\) will be the new \(m\) we shall work with in the coming predictions.

To see that the prediction algorithm as described here, is indeed an **efficient predictor**, we have to prove the following facts summarized in Lemma 3. (Lemma 3 is analogous to Lemma 1 for the second stage).

**Lemma 3:** The following claims hold for a predictor predicting a non-reduced polynomial time \(\Phi\)-generator.

a) The number \(m'\) computed above is a nontrivial divisor of \(m\) and a multiple of the modulus \(m\).
b) The total number of mistakes made by the predictor during the second stage is polynomial in \( \log m \) and \( k \).

c) The prediction time for each \( s_j \) during the second stage is polynomial in \( \log m \) and \( k \).

Proof:

a) Recall that \( m' = \gcd(m, p - s_j) \), so it is a divisor of \( m \). It is a nontrivial divisor because \( p \) and \( s_j \) are reduced mod \( m \) respectively, and then their difference is strictly less than \( m \). It cannot be zero because \( p \neq s_j \), as follows from the incorrectness of the prediction. The proof that \( m' \) is a multiple of \( m \) is similar to that of part a) of Lemma 1. It is sufficient to show that \( p - s_j \) is a multiple of \( m \), since \( m' \) is itself a multiple of \( m \). We show this by proving \( p \equiv s_j \pmod{m} \):

\[
p = s(i-1) \cdot c \pmod{\hat{m}} \quad \text{(by definition of } p )
\]
\[
= \alpha \cdot \overline{B(i-1)} \cdot c \pmod{m} \quad \text{(by (3))}
\]
\[
= \alpha \cdot B_i \pmod{\hat{m}} \quad \text{(c is a solution } \overline{B(i-1)} \cdot x = B_i \pmod{\hat{m}})\)
\[
= s_i \pmod{m} \quad \text{(By definition of } s_i \pmod{2})
\]

As \( m \) divides \( \hat{m} \) part a) follows.

b) The possible mistakes during the second stage are of two types. Mistakes of the first type happen when a solution to the above congruential equations does not exist. This implies the independence modulo the current \( \hat{m} \) of the corresponding \( B_i \). In fact, this \( B_i \) is also independent \( \pmod{\hat{m}_0} \), where \( \hat{m}_0 \) is the modulus computed at the end of the first stage. This follows from the property of being every \( \hat{m} \) a divisor of \( \hat{m}_0 \). By Lemma 2, we have that the number of independent vectors \( \pmod{\hat{m}_0} \) is at most \( k \log \hat{m}_0 \). Therefore the number of mistakes by lack of a solution is bounded by this quantity too. The second type of mistakes is when a solution exists but the computed prediction is incorrect. Such a mistake can occur only once per \( \hat{m} \). After it occurs, a new \( \hat{m} \) is computed. Thus, the total number of such mistakes is as the number of different \( \hat{m} \)'s computed during the algorithm. These \( \hat{m} \)'s form a decreasing sequence of positive integers in which every element is a divisor of the previous one. The first (i.e. largest) element is \( \hat{m}_0 \) and then the length of this sequence is at most \( k \log \hat{m}_0 \). Consequently, the total number of mistakes during the second stage is at most \( (k + 1) \log \hat{m}_0 \), and by Lemma 1 part d) this number is polynomial in \( \log m \) and \( k \).

c) By our assumption of the polynomiality of the functions \( \Phi_j \) when working on the vectors \( s(i) \), it is clear that the computation of each \( B_i \pmod{\hat{m}} \), takes time that is polynomial in \( \log m \) and \( k \). We only need to show that a solution to \( \overline{B(i-1)} \cdot x = B_i \pmod{\hat{m}} \) can be
computed in time polynomial in \( \log m \) and \( k \). A simple method for the solution of a system of linear congruences like the above, is described in [6] (and [3]). This method is based on the computation of the Smith Normal Form of the coefficients matrix in the system. This special matrix and the related transformation matrices, can be computed in polynomial time, using an algorithm of [11]. Thus, finding a solution to the above system (or deciding that no one exists) can be accomplished in time polynomial in \( \log m \) and \( k \). Therefore the whole prediction step is polynomial in these parameters.  

Combining Lemmas 1 and 3 we get

**Theorem:** For every non-reduced polynomial-time \( \Phi \)-generator the predicting algorithm described above is an efficient predictor.

As a special case we get

**Corollary:** Every multivariate polynomial recurrence generator is efficiently predictable.
4. CONCLUDING REMARKS

Our prediction results concern number generators outputting all the bits of the generated numbers, and does not apply to generators that output only parts of the numbers generated. Recent works treat the problem of predicting linear congruential generators which output only parts of the numbers generated [9, 10, 13, 18].

A theorem by Yao [19] states that pseudorandom (bit) generators are unpredictable by polynomial-time means if and only if they pass any polynomial time statistical test. That is, predictability is a universal statistical test in the sense that if a generator is unpredictable, then it will pass any statistical test. Thus, a generator passing this universal test will be suitable for any "polynomially bounded" application. Nevertheless, for specific applications, some weaker generators may suffice. As an example, for their use in some simulation processes, all that is required from the generators is some distribution properties of the numbers generated. In the field of Probabilistic Algorithms the correctness of the algorithm is often analyzed assuming the total randomness of the coin tosses of the algorithm. However, in special cases a more relaxed assumption is possible. For example Bach [2] shows that simple linear congruential generators suffice for guaranteeing the correctness and efficiency of some probabilistic algorithms, even though these generators are clearly predictable. In [7] linear congruential generators are used to "expand randomness". Their method allows the deterministic "expansion" of a truly random string into a sequence of pairwise independent pseudorandom strings.

Provable unpredictable generators exist, assuming the existence of one-way permutations [4, 19]. In particular, assuming the intractability of factoring, an efficient pseudorandom bit generator can be presented [5, 1]. This generator outputs the bit sequence \( b_1, b_2, \ldots \), where \( b_i \) is the least significant bit of \( s_i \), \( s_i \equiv s_{i-1}^2 \pmod{m} \), and \( m \) is the product of two large primes.

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