Decomposing Solution Sets of Polynomial Systems:
A New Parallel Monodromy Breakup Algorithm

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Abstract

We consider the numerical irreducible decomposition of a positive dimensional solution set of a polynomial system into irreducible factors. Path tracking techniques computing loops around singularities connect points on the same irreducible components. The computation of a linear trace for each factor certifies the decomposition. This factorization method exhibits a good practical performance on solution sets of relative high degrees.

Using the same concepts of monodromy and linear trace, we present a new monodromy breakup algorithm. It shows a better performance than the old method which requires construction of permutations of witness points in order to break up the solution set. In contrast, the new algorithm assumes a finer approach allowing us to avoid tracking unnecessary homotopy paths.

As we designed the serial algorithm keeping in mind distributed computing, an additional advantage is that its parallel version can be easily built. Synchronization issues resulted in a performance loss of the straightforward parallel version of the old algorithm. Our parallel implementation of the new approach bypasses these issues, therefore, exhibiting a better performance, especially on solution sets of larger degree.

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1 INTRODUCTION

1.1 Problem Statement

As polynomial equations emerge more and more often in various fields of science and engineering, the question of simplification of polynomials and polynomial systems becomes of the most importance. How can we simplify? One way to understand better the solution set of a polynomial is to factor it; equivalently, in case of a polynomial system we talk about finding an irreducible decomposition of its solution set, a central problem in numerical algebraic geometry \[34, 35\].

A widely known family of polynomial systems used for benchmarking is “cyclic \(n\)-roots”, which arose in Fourier analysis \[3, 4\]. The case \(n = 4\) is our running example:

\[
\begin{align*}
    x_1 + x_2 + x_3 + x_4 &= 0 \\
    x_1x_2 + x_2x_3 + x_3x_4 + x_4x_1 &= 0 \\
    x_1x_2x_3 + x_2x_3x_4 + x_3x_4x_1 + x_4x_1x_2 &= 0 \\
    x_1x_2x_3x_4 - 1 &= 0.
\end{align*}
\]

(1)

This system has a one dimensional solution component of degree four, which becomes obvious by the substitution \(x_3 = -x_1\) and \(x_4 = -x_2\). After this substitution, the first three equation vanish and the last equation simplifies to \(x_1^2x_2^2 - 1 = (x_1x_2 - 1)(x_1x_2 + 1)\), the curve of degree four factors in two irreducible quadrics.

We denote systems of polynomial equations by \(f(x) = (f_1(x), \ldots, f_m(x)) = 0\), where \(f_i \in \mathbb{C}[x] = \mathbb{C}[x_1, \ldots, x_n]\) for all \(i\). Very often, the coefficients are known with limited accuracy. The solution set \(V\) to \(f(x) = 0\) is naturally organized into pure dimensional solution sets \(V = [V_0, V_1, \ldots, V_n]\), where \(\dim(V_k) = k\). A numerical representation of a pure dimensional solution set \(V_k\) is a witness set \[29, 35\], which consists of

1. the polynomials \(f_i\) (\(i = 1, \ldots, m\));

2. \(k\) linear equations \(L(x) = (L_1(x), \ldots, L_k(x)) = 0\) with generic coefficients describing \(k\) generic hyperplane slices;

3. a list \(W\) of \(\deg(V_k)\) solutions to the system \(f(x) = L(x) = 0\).

By the generic choice of the coefficients of the \(L(x) = 0\), the \(k\) hyperplanes defined by \(L\) cut out exactly as many isolated regular solutions on \(V_k\) as \(\deg(V_k)\).

Notice how the treatment of positive dimensional solution sets is reduced to dealing with collections of generic points. Using slack variables we reduce overdetermined polynomial systems to systems with as many variables as unknowns \[28\]. For a system like cyclic 4-roots in \[11\], we add one slack variable \(z\).
to the system:

\[
\begin{align*}
    x_1 + x_2 + x_3 + x_4 + b_1 z &= 0 \\
    x_1 x_2 + x_2 x_3 + x_3 x_4 + x_4 x_1 + b_2 z &= 0 \\
    x_1 x_2 x_3 x_4 + x_3 x_4 x_1 + x_4 x_1 x_2 + b_3 z &= 0 \\
    x_1 x_2 x_3 x_4 - 1 + b_4 z &= 0 \\
    c_0 + c_1 x_1 + c_2 x_2 + c_3 x_3 + c_4 x_4 + z &= 0,
\end{align*}
\]

where the coefficients \(b_1, \ldots, b_4, c_0, c_1, \ldots, c_4\) are randomly chosen complex numbers. The extra linear equation reduces the dimension of the solution set by one and we find generic points on the curve as regular solutions with \(z = 0\). For the cyclic 4-roots system the witness set contains four generic points, as the degree of the curve equals four.

The main question now is: given a positive dimensional solution set \(V\), can we find its decomposition into the irreducible components? In the language of witness sets this interprets as: given a witness set \((f, L, W)\) of \(V\), can we find a decomposition \(W = W^{(1)} \sqcup \ldots \sqcup W^{(r)}\) such that for all \(i\) the witness set \((f, L, W_i^{(i)})\) represents an irreducible component of \(V\)?

Finding polynomial time algorithms for the factorization of multivariate polynomial with approximate coefficients was posed in [20] as one of the challenges in symbolic computation. This challenge received a lot of attention [6, 7, 9, 8, 13, 14, 15, 19, 24, 33]; see [7] for a nice description of recent methods.

1.2 Numerical Homotopies define Loops around Singularities

In [30], a new numerical algorithm using homotopy continuation methods was proposed to decompose a positive dimensional solution set into irreducible factors. Linear traces were proposed in [31] to certify a numerical irreducible decomposition. The implementation [32] was adjusted to the important special case of factoring one single multivariate complex polynomial in [33].

In this section we outline the idea of exploiting monodromy using homotopies to define loops around singularities. Assume two witness sets \((f, L_1, W_1)\) and \((f, L_2, W_2)\) represent the same positive dimensional irreducible component \(V\). Consider the system \(H_{\gamma, L_1, L_2}(x, t)\):

\[
\begin{align*}
    f(x) &= 0; \\
    (1 - t) L_1(x) + \gamma t L_2(x) &= 0. \\
\end{align*}
\]

where \(\gamma\) is a generic nonzero complex number. Then, due to the generic choice of \(\gamma\), for a fixed value of \(t\) the solutions to \(H_{\gamma, L_1, L_2}(x, t)\) are all isolated. In particular, these are \(W_1\) for \(t = 0\) and \(W_2\) for \(t = 1\).

Tracking solutions of \(H_{\gamma, L_1, L_2}(x, t)\) as \(t\) varies from 0 to 1 defines a 1-to-1 map \(\phi_{\gamma, L_1, L_2} : W_1 \rightarrow W_2\). If the composition of two such maps defines a loop around a singularity for some \(t\), then a permutation of the points of a witness set is obtained, in particular:

\[
\pi_{\gamma, L_1, L_2} = \phi_{\gamma, L_2, L_1} \circ \phi_{\gamma, L_1, L_2}
\]
is a permutation of \( W_1 \). All permutations that arise in this fashion form a subgroup of the symmetry group acting on \( W_1 \) and the orbits of this action are witness sets that represent irreducible components.

The idea to exploit monodromy first appeared in a theoretical complexity study \[2\]. Although our approach does not need to know the precise location of the singularities, one could as in \[11\] compute those for algebraic curves, see also the command \texttt{algcurves[monodromy]} in Maple.

The algorithm in \[30\] collects points connected by loops into the same witness sets which converge to numerical representations of the irreducible components. In \[31\], a stop criterion for this algorithm was presented, using the linear trace. We explain this trace test on a system like cyclic 4-roots. Note our program only works with generic points obtained as solutions of \( \mathbf{2} \) and does not have a symbolic polynomial of degree four. Via a generic projection we map the points in 4-space down to the plane. If two of the four points to belong to the same irreducible component, there must exist a quadratic polynomial \( p(x, y) \) vanishing at those two points and at any point of the quadratic irreducible factor. The linear trace is then defined rewriting \( p(x, y) \) as \( p(x, y(x)) \):

\[
p(x, y(x)) = (y - y_1(x))(y - y_2(x))
\]

\[
= y^2 - (y_1(x) + y_2(x))y + y_1(x)y_2(x)
\]

\[
= y^2 - t_1(x)y + t_2(x),
\]

where \( t_1(x) \) is the linear trace. If \( t_1 \) was not linear, then \( \text{deg}(p) > 2 \). So \( t_1(x) = ax + b \), for some \( a \) and \( b \) to be determined by interpolation at \( x = x_0 \) and \( x = x_1 \), with corresponding \( y \)-values in \((x_0, y_{01}), (x_0, y_{02}), (x_1, y_{11})\), and \((x_1, y_{12})\). If for an additional sample, at \( x = x_2 \) with corresponding \( y \)-values \((x_2, y_{21}), (x_2, y_{22})\), we have \( t_1(x_2) = y_{21} + y_{22} \), then we have an irreducible quadratic factor, otherwise the two points do not lie on the same factor.

The linear trace test, called zero-sum relations, was first introduced in \[27\] and further developed in \[25, 26\]. For factors of small to moderate degree, the linear trace test can be applied in an exhaustive combinatorial enumeration as was proposed in \[13, 14, 23\], \[24\] and improved in \[0\].

\subsection*{1.3 Parallel Algorithms}

Homotopy continuation methods are very well suited for parallel processing as after distributing the path tracking jobs among the computers in the network, no further communications are needed, see \[115, 17\] for granularity issues. For computational algebraic geometry, this implies that homotopy methods can solve much larger polynomial systems than methods in computer algebra which are harder to adapt to parallel computers \[21\]. One recent example is the solution of the cyclic 13-roots problems with PHoM \[10, 16\] for which 2,704,156 paths were tracked.

Modern homotopies in numerical algebraic geometry often appear in a sequence like the Pieri homotopies \[37\] where the start solutions of one homotopy lie at the end of paths defined by another homotopy. The homotopies to factor
positive dimensional solution sets raise job scheduling issues as the decision to
certain track paths depends on the outcome of other paths. This paper can be
regarded as a solution to the job scheduling problems raised in \[22\].

The parallel algorithm proposed in \[22\] exhibited a good speedup in the path
tracking jobs, but the certification with linear traces executed in only by the
master node before the scheduling of new path tracking jobs diminished the
overall performance as all nodes were idling waiting for the assignment of new
path tracking jobs. At the end of \[22\] we outlined a probabilistic complexity
study simulated in Maple, suggesting various job scheduling techniques. In this
paper we report on its parallel implementation confirming the efficiency of the
approach.

2 USING MONODROMY MORE EFFICIENTLY

The monodromy breakup algorithm of \[30, 31\] is sketched by Algorithm \[2.1\].

On input is a witness set \(W_L\) and on output a partition of \(W_L\), corresponding
to the irreducible decomposition.

Algorithm 2.1 Monodromy Breakup certified by Linear Trace:

\[ \mathcal{P} = \text{Breakup}(W_L, d, N) \]

\[ \text{Input: } W_L, d, N. \]
\[ \text{Output: } \mathcal{P}. \]

0. initialize \(\mathcal{P}\) with \(d\) singletons;
1. generate two slices \(L'\) and \(L''\) parallel to \(L\);
2. track \(d\) paths for witness set with \(L'\);
3. track \(d\) paths for witness set with \(L''\);
4. for \(k\) from 1 to \(N\) do
   4.1 generate new slices \(K\) and a random \(\gamma\);
   4.2 track \(d\) paths to a new slice;
   4.3 generate a new random \(\gamma\);
   4.4 track \(d\) paths to return to the base slice;
   4.5 compute the permutation and update \(\mathcal{P}\);
   4.6 if linear trace test certifies \(\mathcal{P}\)
      then leave the loop;
   end if;
end for.

Our first parallel implementation of this algorithm, described in \[22\], uses a
master/servant model where the master node distributes the paths among the
available processors in the network. According to our experimental results, a
sizeable speedup is achieved by distributing the routine path-tracking jobs to
different nodes. However, a probabilistic study in \[22\] suggested that we can
save some work by taking a smaller one-path-one-point tracking job as an atomic
task.

Following the previous discussion, we take two generic slices \(L_1\) and \(L_2\) (in
this case these are hyperplanes) and look at the witness points \(W_1\) and \(W_2\).
Consider the bipartite graph with vertices $W_1$ on one side and $W_2$ on the other.

One atomic step of the monodromy breakup algorithm consists of creating a map like $\phi_{\gamma, L_1, L_2}$. We can visualize such a map by connecting the points of $W_1$ and $W_2$ that map into each other with an edge.

As you may see in our example, in order to create one permutation we need to construct 8 edges in the graph. If one is lucky then it may take [MBA-P] only one permutation to decompose cyclic4:

The connected components of the graph in Figure 2 correspond to the two witness sets that, in turn, represent two irreducible components of the solution set of cyclic4: two quadric curves.

In fact 2 out of 8 edges in Figure 2 can be removed keeping the connected components still connected, see Figure 2. Since each edge can be created by tracking only one point of a witness set, we may avoid doing extra work by trying to create as few edges as possible.
3 A NEW ALGORITHM

In this section we first describe the sequential version of our new monodromy breakup algorithm before addressing its parallel execution.

3.1 Serial Version

The flow chart of our new algorithm is show in Figure 4. As in Algorithm 2.1, we also have the initialization of the “trace grid”, which are the two witness sets on two parallel slices needed to certify the irreducible decomposition using linear traces. While Algorithm 2.1 first completes all the loops for all points in a witness set before proceeding to the next level, our new approach initializes $s$ new witness sets which are available for generating loops.

The main loop of the new algorithm shown in Figure 4 leaves much freedom to complete loops between any two slices. For every slice, the algorithm keeps track of the number of loops that did not yield a permutation, stored in $N_{rej}$. Based on these statistics, the algorithm can discriminate against slices which were not productive in the past and select those slices which led to more new permutations.

As explained in the previous section, the algorithm typically returns with a certified decomposition before all loops are completed. This is the main reason why on single processors, our new algorithm outperforms Algorithm 2.1. At the same time, the new algorithm is more suitable for parallel execution, as we will explain next.

3.2 Parallel Version

The parallel version of our new algorithm runs in a master/servant model. The initialization phase is very similar to the initialization of the parallel version of Algorithm 2.1 with the master distributing path tracking jobs evenly among all nodes. After the initialization, the master node keeps looking for available
path tracking nodes to assign paths and the other nodes are either busy tracking paths or ready to start new path tracking jobs.

Compared to our previous parallel algorithm described in [22], the computation of the linear trace by the master node is now interleaved by path tracking jobs performed on the other nodes.

4 EXPERIMENTAL RESULTS

Our algorithms are implemented using the path tracking routines in PHCpack [36], extended in [32] with facilities for a numerical irreducible decomposition. As in [37], we apply MPI for message passing. Our main program is written in C and links with the interface of PHCpack.

Our equipment consists of two personal cluster machines purchased from Rocketcalc (www.rocketcalc.com) for a total of 12 2.4Ghz CPUs, served by a Dell workstation with two dual 2.4Ghz processors. So in total we have 14 processors at our disposal.

4.1 Plain Parallel Path Tracking

In Table 1 we show with three runs the main defect of our first parallel implementation presented in [22]. While we have no real control over the number of loops it takes to complete the factorization, we observe from the data in Table 1 that as the total number of loops increases, the work done by the master node increases.

| #loops | 4        | 6        | 9        |
|--------|----------|----------|----------|
| min track | 8.0 sec  | 10.9 sec | 18.5 sec |
| max track | 10.8 sec | 15.7 sec | 21.8 sec |
| master   | 1.8 sec  | 3.8 sec  | 7.6 sec  |
| total    | 12.6 sec | 19.5 sec | 29.4 sec |

Table 1: Three runs with the first parallel monodromy breakup algorithm, executing respectively 4, 6, and 9 loops to factor a curve of degree 144 in 8 space on 14 processors. We report the minimal and maximal time the nodes spent tracking paths, and the time spent by the master node certifying the decomposition and scheduling the jobs.

Although the cyclic 8-roots system is a problem of modest size, we already observe in Table 1 that for 9 loops, more than 25% of the time is spent by the master node, while all the other nodes are idling. For larger problems and more processors, the poor performance of this first implementation will become even more apparent.
4.2 Performance of the new Algorithm

Table 2 reports on five runs done on 14 processors to decompose the curve of degree 144 defined by the cyclic 8-roots system. For three of the five runs we used three new slices. In the last two runs we see that the total time decreases if we use only two new slices.

| #runs | 3 new slices | 2 new slices |
|-------|--------------|--------------|
|       |              |              |
| initial | 8.73 | 9.01 | 8.89 | 6.54 | 6.98 |
| master  | 6.06 | 6.22 | 6.18 | 6.67 | 7.10 |
| min track | 5.96 | 6.16 | 6.07 | 6.60 | 7.02 |
| max track | 6.06 | 6.24 | 6.23 | 6.11 | 7.15 |
| total   | 14.9 | 15.4 | 15.3 | 13.4 | 14.2 |

Table 2: Five runs with our new parallel monodromy breakup algorithm, three times with 3 new slices and two times with 2 new slices. We report the time used for initialization, the time spent by the master node, the minimal and maximal time for the nodes spent tracking paths, and the total time. All reported times are expressed in seconds.

In Table 2 we see an even distribution of the time spent by the nodes. Using fewer slices reduces the initialization time at the expense of a slightly higher running time in the main loop.

Comparing to the timings in Table 1 we do not notice such a wide fluctuation in the total execution time between different numbers of loops. The total execution time of the most favorable situation reported in Table 1 is only slightly lower than the best total time in Table 2.

Finally, we report on a calculation of a larger example, the ideal of adjacent 2-by-2 minors of a general 2-by-9 matrix of 18 unknowns, see [12] and [18]. This system in 18 variables defines a 10-dimensional surface of degree 256 which factors in 34 irreducible components. The total execution time of our new monodromy breakup algorithm on 14 processors is 97.1 seconds, of which 62.7 are spent on the initialization, 33.8 seconds by the master node in the main loop while the time path tracking on the other nodes fluctuated between 32.9 and 35.6 seconds.

The performance of our first parallel algorithm on this system is even more erratic. The very best complete run of 3 monodromy loops took 122.9 seconds on 14 processors, where the path tracking time ranged between 75.9 and 104.4 seconds. Even on this very best run, our new algorithm still takes only 80% of the time spent by the first parallel monodromy breakup algorithm of [22].
5 CONCLUSIONS

In this paper we report on the development and performance of a new monodromy breakup algorithm. Experimental results show a more predictable and regular performance than our first parallel implementation of [22]. Thanks to this increased performance, it is now possible to factor solution sets of larger degrees.

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Initialization:
P := \{ \{a\} \mid 1 \leq a \leq d, \{a\} \text{ is not a component}\}
Q := \{\{a\} \mid 1 \leq a \leq d, \{a\} \text{ is a component}\}
N_{tot} = N_{red} = 0

Construct \( s \) witness sets using random slices
Label the witness points on the slices
Construct the trace grid

\#P ≠ 0

yes

Pick the smallest part \( p \in P \) and a label \( a \in p \)

Pick two slices \( L_1 \) and \( L_2 \)

\( b := \text{track}(a, L_1, L_2) \)
\( N_{tot} := N_{tot} + 1 \)
Find \( q \in P \) that contains the label \( b \)

\( q = p \)

yes

\( N_{rej} := N_{rej} + 1 \)

Return \( Q, N_{tot}, N_{rej} \)

Merge \( p \) and \( q \):
\( P := P \setminus \{p, q\} \)
\( P := P \cup \{p \cup q\} \)

Is \( p \cup q \) irreducible?

yes

\( P := P \setminus \{p \cup q\} \)
\( Q := Q \cup \{p \cup q\} \)

\( P := P \setminus \{p, q\} \)
\( Q := Q \cup \{p \cup q\} \)

Figure 4: The flow chart of our new Monodromy Breakup Algorithm