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To cite this version:
Sylvain Collange, Mioara Joldes, Jean-Michel Muller, Valentina Popescu. Parallel floating-point expansions for extended-precision GPU computations. The 27th Annual IEEE International Conference on Application-specific Systems, Architectures and Processors (ASAP), Jul 2016, London, United Kingdom. hal-01298206v2

HAL Id: hal-01298206
https://hal.archives-ouvertes.fr/hal-01298206v2
Submitted on 11 Apr 2016

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Parallel floating-point expansions for extended-precision GPU computations

Sylvain Collange, Mioara Joldes, Jean-Michel Muller and Valentina Popescu

Abstract—GPUs are an important hardware development platform for problems where massive parallel computations are needed. Many of these problems require a higher precision than the standard double floating-point (FP) available. One common way of extending the precision is the multiple-component approach, in which real numbers are represented as the unevaled sum of several standard machine precision FP numbers. This representation is called a FP expansion and it offers the simplicity of using directly available and highly optimized FP operations.

In this article we present new data-parallel algorithms for adding and multiplying FP expansions specially designed for extended precision computations on GPUs. These are generalized algorithms that can manipulate FP expansions of different sizes (from double-double up to a few tens of doubles) and ensure a certain worst case error bound on the results.

Index Terms—floating-point arithmetic, floating-point expansions, high precision arithmetic, multiple-precision arithmetic, graphics processing unit, parallel computations

1 INTRODUCTION

In numerical computing, we sometimes encounter calculations that require more precision than the one offered by current processors. A way of handling such higher-precision calculations is to represent real numbers by floating-point (FP) expansions, i.e., by nonevaluated sums of FP numbers. Several slightly different definitions of what a FP expansion is have been introduced in the literature [1], [2]. Also, several different arithmetic algorithms for manipulating them have been suggested. Choosing between these algorithms frequently depends on a compromise between accuracy, speed, and safety (because some of the more complex algorithms are just heuristic: they do not come with a proof). With Graphics Processing Units (GPUs) oriented implementation in mind, other important aspects are parallelism and locality. Should we try to parallelize the arithmetic algorithms that manipulate FP expansions (or, rather, build variants that are suitable for parallelization), or should we keep them sequential and try to parallelize at a higher level? In a previous study [3], some of us have dealt with an “embarrassingly parallel” problem with compact intermediate data. However, many applications do not provide as much parallelism. Even for those that do, locality can be a problem. Increasing precision of sequential arithmetic operations requires a corresponding increase in the amount of intermediate data to keep. Thus, parallel arithmetic algorithms are attractive not just by the extra parallelism they provide, but also by the locality improvements they enable. Here, with different numerical problems in mind, we parallelize the addition and multiplication of FP expansions on GPUs.

Section 2 recalls the basic notions related to FP expansions. In Sections 3 and 4 we present data-parallel algorithms for addition and multiplication, respectively, of FP expansions. The implementation details are given in Section 5 followed by some performance assessment in Section 6. We finish by concluding our work in Section 7.

2 FLOATING-POINT EXPANSIONS

A normal binary precision-$p$ floating-point (FP) number is a number of the form

$$x = M_x \cdot 2^{e_x - p + 1},$$

with $2^{p-1} \leq |M_x| \leq 2^p - 1$. The integer $e_x$ is called the exponent of $x$, and $M_x \cdot 2^{e_x - p + 1}$ is called the significand of $x$. We denote accordingly to Goldberg’s definition $\text{ulp}(x) = 2^{e_x - p + 1}$ [4, Chap. 2].

A natural extension of the notion of double-double or quad-double is the notion of floating-point expansion.

Definition 2.1. A FP expansion $u$ with $n$ terms is the unevaluated sum of $n$ FP numbers $u_0, u_1, \ldots, u_{n-1}$, in which all nonzero terms are ordered by magnitude (i.e., if $v$ is the sequence obtained by removing all zeros in the sequence $u$, and if sequence $v$ contains $m$ terms, $|v_i| \geq |v_{i+1}|$, for all $0 \leq i < m-1$).

Arithmetics on FP expansions have been introduced by Priest [1], and later on by Shewchuk [2].

To make sure that such an expansion carries significantly more information than only one FP number, it is required that the $u_i$’s do not “overlap”. The notion of (non-)overlapping varies depending on the authors. We give here the definition used by Priest in his initial work and we introduce a new one, with a weaker condition, that allows for a more relaxed handling of the FP expansions.
We specify first that even if a FP expansion may contain interleaving zeros, all the definitions that follow apply only to the non-zero terms of the expansion (i.e., the array $v$ in Definition 2.1).

**Definition 2.2.** Assuming $x$ and $y$ are normal numbers with representations $M_x \cdot 2^{e_x-2^p+1}$ and $M_y \cdot 2^{e_y-2^p+1}$ (with $2^{p-1} \leq |M_x|, |M_y| \leq 2^p - 1$), they are $\mathcal{P}$-nonoverlapping (that is, nonoverlapping according to Priest’s definition [5]) if $|e_y - e_x| \geq p$.

**Definition 2.3.** An expansion is $\mathcal{P}$-nonoverlapping (that is, nonoverlapping according to Priest’s definition [5]) if all its components are mutually $\mathcal{P}$-nonoverlapping.

Intuitively, the stronger the sense of the nonoverlapping definition, the more difficult it is to obtain, implying extra manipulation of the FP expansions. In order to save operations we chose to use a slightly weaker sense of nonoverlapping, referred to as ulp-nonoverlapping, that we define in what follows.

**Definition 2.4.** An expansion $u_0, u_1, \ldots, u_{n-1}$ is ulp-nonoverlapping if for all $0 < i < n$, we have $|u_i| \leq \text{ulp}(u_{i-1})$.

In other words, the components are either $\mathcal{P}$-nonoverlapping or they overlap by one bit, in which case the second component is a power of two.

**Remark 2.5.** Note that for $\mathcal{P}$-nonoverlapping expansions we have $|u_i| \leq \frac{2^{p-1}}{2^i} \text{ulp}(u_{i-1})$.

Depending on the nonoverlapping type of an expansion, when using standard FP formats for representation, the exponent range forces a constraint on the number of terms. The largest expansion can be obtained when the largest term is close to overflow and the smallest is close to underflow. We remark that, when using any of the above nonoverlapping definitions, for the two most common FP formats, the constraints are:

- for double-precision (exponent range $[-1022, 1023]$) the maximum expansion size is 39;
- for single-precision (exponent range $[-126, 127]$) the maximum is 12.

In this article we deal with so called parallel FP expansion, i.e., the expansion is stored on parallel execution threads, with one term/thread. This implies that the user has to launch as many threads as the expansion size.

The majority of algorithms performing arithmetic operations on FP expansions are based on the so-called error-free transforms (EFT), such as the algorithms $2\text{Sum}$, Fast$2\text{Sum}$, Dekker’s product and $2\text{MultFMA}$ (presented for instance in [4]), that make it possible to compute both the result and the error of a FP addition or multiplication. This implies that each such EFT applied to two FP numbers, returns still two FP numbers. These EFT can be extended to work on several inputs by chaining, resulting in the so-called distillation algorithms [6], for summing several FP numbers. From these we make use of the $\text{VecSum}$ algorithm [2], [7], which is simply a chain of $2\text{Sum}$ that performs an EFT on $n$ FP numbers.

A potential problem appears when subsequent computations are done using these results; the size of the exact result is going to increase more and more. To avoid this, some “truncation” methods (both on-the-fly or a-posteriori) may be used to compute only an approximation of the exact result. Also, so-called (re-)normalization algorithms are used to render the result nonoverlapping, which implies also a potential reduction in the number of components.

### 3 Data-parallel addition algorithm for FP expansions

An algorithm that performs addition of two FP expansions $x$ and $y$ with $n$ and $m$ terms, respectively, will return a FP expansion with at most $n + m$ terms as the exact result. This implies a continuous increase in the number of terms as subsequent computations are done using the obtained result. This is why, in practice, the results are “truncated” (either on-the-fly or a-posteriori) to obtain only an approximation of $x + y$.

Many variants of algorithms that compute the sum of two FP expansions have been presented in the literature [1], [2], [8], [6], each using different methods and having different complexities, but, from our knowledge, none of these algorithms are implemented in parallel and, even more, some of them are highly sequential, making them unsuitable for parallel architectures.

In what follows we will present a safe data-parallel algorithm for adding FP expansions that offers a tight error bound on the result and it allows to prove a clear constraint between the terms of the result. We also present a fast version with more relaxed error bounds based on the same scheme that may be used if computations are not cancellation-prone.

The safe data-parallel summation algorithm is presented in Fig. 1 and Algorithm 1. All arithmetic operations including EFTs like $2\text{Sum}$ are performed in parallel element-wise on $R$-element vectors. We assume vectors can be merged and elements inside a vector can be shuffled. These assumptions make the algorithms applicable to most SIMD units, including Intel SSE/AVX instruction set extensions [9] and recent Nvidia GPUs [10].

For the sake of simplicity, we only present here the “input-$R$-output-$R$” version of the algorithm, even though the generalized version allows for different input sizes.

The algorithm is based on a pipelined error propagation. We start by adding the first elements of each expansions, $x_0$ and $y_0$ on the first vector component. We continue to add the rest of the elements on the first component one by one and propagating the error upwards, to the other vector components. When we run out of elements to add we continue to propagate the errors for another $R - 1$ steps by injecting $0$s on the first component. In the last step of the algorithm there is no need to use EFT, since we are not going to propagate the errors anymore; this is why we use only simple addition.
It is easily seen that the parallel scheme presented in Fig. 1 can be reduced to a sequential one, that looks like this:

![Sequential Scheme](image-url)

If there is a “Sterbenz relation” between $x_0$ and $y_0$ (i.e., if they are of opposite signs and $|\frac{x_0}{y_0}| \leq |y_0| \leq |2x_0|$) then $e_{01} = 0$ and $s_{01} = x_0 + y_0$. This implies that $s_{12} = e_{02}$ and $e_{12} = 0$, and so on.

In this case we end up propagating a 0, to the end of the result expansion, and we are left with the same scheme as we began with. This means that we can eliminate the case in which we have a “Sterbenz relation”.

Now let us prove the following proposition that refers to only one horizontal line of the scheme (Fig. 2).

![Simple Scheme](image-url)

**Proposition 3.1.** Let us assume $a = a_0, a_1, \ldots, a_{n-1}$, an array of FP numbers that satisfy $|a_i| \leq 2^{-i(p-1)+\delta}|a_0|$, for some (presumably small) integer $\delta$ and $|t| \leq 2^{-p+\ell}|a_0|$, for
some (presumably small) integer \( \ell \). If \( s = s_0, s_1, \ldots, s_n \) is the sum obtained by adding \( t \) to \( a \) as shown in Fig. 2 (from left to right, propagating the error), then all the terms in \( s \) satisfy: \( |s_i| \leq 2^{-i(p-1)+\delta+1} \), for all \( 0 \leq i \leq n \).

**Proof:** From the proof of the algorithm 2Sum we know that \( |e_i| \leq \frac{1}{2} \ulp(s_i) = 2^{-p} |s_i| \).

From
\[
|s_0|(1 - 2^{-p}) \leq |a_0 + t| \leq |s_0|(1 + 2^{-p}) \quad \text{and} \quad |a_0|(1 - 2^{-p+\ell}) \leq |a_0| |(1 + 2^{-p+\ell}),
\]
we get
\[
|s_0| \frac{1 - 2^{-p}}{1 + 2^{-p+\ell}} \leq |a_0| \leq |s_0| \frac{1 + 2^{-p}}{1 - 2^{-p+\ell}}.
\]

(1)

It follows that
\[
|a_1| \leq 2^{-(p-1)+\delta} |a_0| \leq 2^{-(p-1)+\delta} \cdot 1 + 2^{-p+\ell} \cdot |s_0|.
\]

This gives
\[
|e_0 + a_1| \leq 2^{-p} \left( 1 + \frac{2^{\delta+1}(1 + 2^{-p})}{1 - 2^{-p+\ell}} \right) |s_0|.
\]

From which we deduce
\[
|s_1| \leq 2^{-p} (1 + 2^{-p}) \left( 1 + \frac{2^{\delta+1}(1 + 2^{-p})}{1 - 2^{-p+\ell}} \right) |s_0|.
\]

(2)

We can continue, by noticing that \( |e_1| \) is bounded by \( 2^{-p} |s_1| \), and bounding \( |a_2| \) by \( 2^{-2(p-1)+\delta} \frac{1 + 2^{-p}}{1 - 2^{-p+\ell}} \cdot |s_0| \).

This gives a bound on \( |e_1 + a_2| \), and a bound on \( s_2 \) is obtained by multiplying that last bound by \( (1 + 2^{-p}) \). An easy induction finally gives
\[
|s_i| < 2^{-i(p-1)+\delta} |s_0|,
\]
with
\[
\theta_i = (1 + 2^{-p})^i + \frac{1}{1 - 2^{-p+\ell}} \sum_{j=1}^{i} 2^{2^{i}(1 + 2^{-p})}{i,j+2}. \quad (4)
\]

One easily finds
\[
\theta_i = (1 + 2^{-p})^i + \frac{2^{i+1}(1 + 2^{-p})}{1 - 2^{-p+\ell}} \left[ \frac{(2^i - (1 + 2^{-p}))^i}{1 - 2^{-p}} \right],
\]

hence,
\[
\theta_i = 2^{i+\delta+1} H_i,
\]
with
\[
H_i = \frac{(1 + 2^{-p})^i}{2^{i+\delta+1}} + \frac{(1 + 2^{-p})^2}{1 - 2^{-p+\ell}} \left( 1 - \frac{2^{i+1}(1 + 2^{-p})^2}{1 - 2^{-p}} \right).
\]

Denote \( u = 2^{-p} \). In all practical cases \( \ell \geq 2 \) and \( \delta \geq 0 \), so that \( H_i \leq G_i \), with
\[
G_i = \frac{1}{2} \left( \frac{1 + u}{2} \right)^i + \frac{(1 + u)^2}{1 - 4u} \left( 1 - \frac{(1 + u)^2}{1 - u} \right).
\]

We have,
\[
G_i = 1 - \frac{1}{2} \left( \frac{1 + u}{2} \right)^i + \frac{u(u+6) - 1 - u}{2u(7-3u)} \left( \frac{1 + u}{1 - 4u} \right).
\]

The only positive term (after the initial “1”) in that sum is \( \frac{u(u+6)}{1 - 4u} \), which is less than \( 7u \) for all pertinent values of \( u = 2^{-p} \). Hence \( G_i \) is less as soon as \( 2^{i+1} \leq 2^p/7 \), which occurs in all practical cases. This gives
\[
|s_i| < 2^{-i(p-1)+\delta} |s_0|,
\]
with \( \delta' = \delta + 1 \).

This concludes our proof.

Hence, in the array of Fig. 1, \( \delta \) is increased by 1 at each line. For instance, if we add two order- \( n \) expansions (i.e., we use \( 2n-1 \) lines in the Array of Fig. 1), then the terms \( s_0, s_1, \ldots \) of the result satisfy
\[
|s_i| \leq 2^{-(p-1)+2n-1} |s_0|,
\]
and the expansion obtained by keeping the first \( n \) terms only represents the sum with an error less than
\[
(2^{-np+3n-1} + 2^{-(n+1)p+3n} + 2^{-(n+2)p+3n+1} + \cdots) \cdot |a_0|,
\]
i.e., with a relative error bounded by a value slightly larger than \( 2^{-np+3n-1} \).

We would like to also mention here a faster, relaxed version of the above algorithm, that requires at most \( R-1 \) steps in order to compute the result (the last step using only simple addition). This algorithm (Fig. 3) offers a worse error bound and it does not ensure the correct result when cancellation occurs, if no re-normalization algorithm is applied on the result. We recommend using this algorithm only for the average case, in which the user is sure that no cancellation may occur.

![Fig. 3: Fast data-parallel FP expansion addition algorithm](image)

### 4 DATA-PARALLEL MULTIPLICATION ALGORITHM FOR FP EXPANSIONS

In general, an algorithm that performs the multiplication of two expansions \( x \) and \( y \) with \( n \) and \( m \) terms, respectively, will return a FP expansion with at most \( 2nm \) terms [1], which, as in the case of addition, implies an increase in the number of terms.
The algorithm that we present (Algorithm 2) computes an approximation of \( x \cdot y \), where \( x \) and \( y \) are two parallel expansion. Here we also present just the “input-R-output-R” version.

**Algorithm 2** Data-parallel algorithm for multiplying FP expansions.

**Input:** FP expansion vectors \( x = (x_0, x_1, \ldots, x_{R-1}) \) and \( y = (y_0, y_1, \ldots, y_{R-1}) \).

**Output:** FP expansion vector \( \pi = (\pi_0, \pi_1, \ldots, \pi_{R-1}) \).

\[
\begin{align*}
1 & : s \leftarrow (0, \ldots, 0) \\
2 & : \pi \leftarrow (0, \ldots, 0) \\
3 & : \text{for } i \leftarrow 0 \text{ to } R - 2 \text{ do} \\
4 & : y' \leftarrow (y_i, y_{i+1}, \ldots, y_{i+(R-2)}) \text{ //Broadcast} \\
5 & : (p, e) \leftarrow 2\text{Prod}(x, y') \\
6 & : (s, e') \leftarrow 2\text{Sum}(s, p) \\
7 & : \pi_1 \leftarrow s_0 \text{//Insert into vector} \\
8 & : s \leftarrow (s_1, s_2, \ldots, s_{R-1}, 0) \text{//Shift left} \\
9 & : \text{while } e \neq 0 \text{ do} \\
10 & : (s, e) \leftarrow 2\text{Sum}(s, e) \\
11 & : e \leftarrow (0, e_0, e_1, \ldots, e_{R-2}) \text{//Shift right} \\
12 & : \text{end while} \\
13 & : \text{while } e' \neq 0 \text{ do} \\
14 & : (s, e') \leftarrow 2\text{Sum}(s, e') \\
15 & : e' \leftarrow (0, e'_0, e'_1, \ldots, e'_{R-2}) \text{//Shift right} \\
16 & : \text{end while} \\
17 & : \text{end for} \\
18 & : p \leftarrow x \cdot y \\
19 & : s \leftarrow s + p \\
20 & : \pi_{R-1} \leftarrow s_0 \text{//Insert into vector} \\
21 & : \text{return } \pi.
\end{align*}
\]

We consider two parallel FP expansions \( x \) and \( y \), each with \( R \) terms and we compute the \( R \) most significant FP components of the product \( \pi = x \cdot y \). We use the following intuition: let \( \varepsilon = \frac{1}{2} \text{ulp}(\pi_0) \), then roughly speaking, if \( \pi_0 \) is of order of \( O(\Lambda) \), then \( e_0 \) is of order \( O(\varepsilon \Lambda) \). This means that for each product \( (p, e) = 2\text{ProdFMA}(x_i, y_j) \), \( p \) is of order \( O(\varepsilon^2 \Lambda) \) and \( e \) of order \( O(\varepsilon^{1+j} \Lambda) \). We truncate the result on-the-fly, by considering only the terms for which \( 0 \leq i + j \leq R - 1 \), since the smaller terms have an order of magnitude much smaller and usually they will not influence the result.

The multiplication algorithm runs as follows: at each iteration \( i \) of the for loop (lines 3-16) we compute \( p + e = x \cdot y \); we add \( p \) to the result of the same order, using an EFT, which also generates an error, \( e' \). After that, using the two while loops (lines 8-11 and 12-15) we propagate the two generated errors, \( e \) and \( e' \) to the lower order results. In the last step of the algorithm, we do not use any EFT, because the errors that are supposed to be computed are going to be of order \( O(\varepsilon R \Lambda) \), and we do not need to propagate them anymore.

This algorithm has the same behavior as the sequential algorithm presented in Fig. 4, but a graphical representation of the parallel one it would be too difficult to read.

While the exact product \( xy \) of two FP expansions \( x \) and \( y \) is computed as

\[
\sum_{i=0}^{R-1} \sum_{j=0}^{R-2} x_i y_j = \sum_{k=0}^{R-2} \sum_{i+j=k} x_i y_j,
\]

in this algorithm we “truncate” it on-the-fly by computing and adding only the relevant part of the scalar products (the first \( \sum_{k=1} k \) individual products) and after that outputting the \( R \) most significant components in the result. In what follows we compute the error bound in two steps: first we compute the error generated by “truncating” the partial products, and second we compute the error given by the discarded errors.

**Proposition 4.1. (Error bound on the truncated products)** Let \( x \) and \( y \) be two ulp-nonoverlapping FP expansions, with \( R \) terms. If, when computing the product \( xy \) we “truncate” the operations by computing and adding only the most significant individual products, the first \( \sum_{k=1} k \) products, then the maximum error that we can obtain is:

\[
\sum_{k=R-2}^{R-1} \sum_{i+j=k} x_i y_j \leq |x_0 y_0| 2^{-(p-1)R} \frac{R-1}{2^{-(p-1)}}.
\]

**Proof:** The maximum error given by the discarded products satisfies:

\[
\sum_{k=R}^{R-2} \sum_{i+j=k} x_i y_j \leq \sum_{k=R}^{R-2} \sum_{i+j=k} 2^{-p(i+j)+i+j} |x_0 y_0| = |x_0 y_0| \sum_{k=R}^{R-2} (2R - 1 - k) 2^{-(p-1)k};
\]

\[
\leq |x_0 y_0| 2^{-(p-1)R} \sum_{k'=0}^{R-2} (R - 1 - k') 2^{-(p-1)k'}.
\]

Fig. 4: Graphical representation of the sequential algorithm that behaves like Algorithm 2.
We now consider the function $\phi(\alpha) = \sum_{k=0}^{\infty} (R-1-k)\alpha^k$ and we get:

$$
\phi(\alpha) = \sum_{k=0}^{\infty} -k\alpha^k + \sum_{k=0}^{\infty} (R-1)\alpha^k;
= -\alpha \frac{d}{d\alpha} \left( \sum_{k=1}^{\infty} \alpha^k \right) + (R-1) \frac{1}{1-\alpha};
= \frac{-\alpha}{(1-\alpha)^2} + \frac{R-1}{1-\alpha}.
$$

Using $\phi(2^{-(p-1)})$, we obtain:

$$
\sum_{k=R}^{2R-2} \sum_{i+j=k} x_i y_j \leq |x_0 y_0| 2^{-(p-1)R} \left( \frac{R-1}{1-2^{-(p-1)}} \right),
$$

and since $\frac{1}{1-2^{-(p-1)}}$ is negative and very small we conclude our proof by:

$$
\sum_{k=R}^{2R-2} \sum_{i+j=k} x_i y_j \leq |x_0 y_0| 2^{-(p-1)R} \frac{R-1}{1-2^{-(p-1)}}. \tag{5}
$$

\[\square\]

**Proposition 4.2.** (Final error bound) Let $x$ and $y$, two ulp-nonoverlapping FP expansions, with $R$ terms. When computing $\pi$, an approximation of $xy$ to $R$ terms, as shown in Algorithm 2 and Fig. 4, the result satisfies:

$$
|xy - \pi| \leq |x_0 y_0| 2^{-(p-1)R} (R-1) \left( 1 + 2^{p-2}(1 + 2^{-p}) + (R^3 - R)((R - 1)!)^2 \right). \tag{6}
$$

**Proof:** From the definition we know that $|x_i| \leq 2^{-(p-1)}|x_0|$ and $|y_j| \leq 2^{-(p-1)}|y_0|$, so we can deduce

$$
|x_i y_j| \leq 2^{-(p-1)(i+j)} |x_0 y_0|.
$$

For computing $\pi_0$ we use only 2MultFMA($x_0, y_0$), and we get $|\pi_0| \leq |x_0 y_0| (1 + 2^{-p})$.

For computing $\pi_1$ we use VecSum#1, with three entries of order $O(\varepsilon^2 \Lambda)$: the error from the previous step, and two partial products, which are less than $2^{-(p-1)} |x_0 y_0|(1 + 2^{-p})$. It is easily seen that all these entries are bounded by the same value. We define the following notation:

$$
\Omega_1 = 2^{-(p-1)} |x_0 y_0|(1 + 2^{-p}). \tag{7}
$$

It follows that $\pi_1 < 3 \cdot 2^{-(p-1)} |x_0 y_0|(1 + 2^{-p})^3$ and the out putted errors are less than $\Omega_2 = 3 \cdot 2^{-(p-1)} |x_0 y_0|(1 + 2^{-p})^3$.

For computing $\pi_2$ we use VecSum#2 which is going to have seven entries of order $O(\varepsilon^2 \Lambda)$: two errors outputted by VecSum#1, bounded by $\Omega_2$; two errors from the previous step’s partial products, which are less than $2^{-(p+1)} |x_0 y_0|(1 + 2^{-p})$; and 3 partial products, less than $2^{-(p-1)} |x_0 y_0|(1 + 2^{-p})$. We observe once more that all the entries are less than $\Omega_2$.

For the induction step we consider VecSum#i - 1. For computing $\pi_i - 1$ we have $(i - 1)^2 + i$ entries, which we assume are all less than $\Omega_{i-1}$. It follows:

$$
\pi_{i-1} < (2\Omega_{i-1}(1 + 2^{-p}) + \Omega_{i-1})(1 + 2^{-p}) + \ldots < (i^2 - i + 1)\Omega_{i-1}(1 + 2^{-p})^{i-1}
$$

and also, the largest error term outputted and implicitly all the others are less than $\frac{1}{2} \ulp(\pi_{i-1})$.

This implies that all error terms are less than:

$$
\begin{align*}
   & (i^2 - i + 1)\Omega_{i-1}(1 + 2^{-p})^{1+2+\ldots+(i^2-i)} \\
   & \leq 2^{-(i+1)p-2} |x_0 y_0| (1 + 2^{-(p-1)})(1 - 2^{-(p-1)}) = \frac{1}{2} \ulp(\pi_{i-1}).
\end{align*}
$$

\[\square\]

Unfortunately, for the multiplication algorithm we are unable to prove any constraints on the terms of the result. Even though cancellation cannot happen when multiplying two FP numbers, it may happen during the summation process, in which case we can get $|\pi_i| < |\pi_j|$, with $i < j$. If this happens we can apply a re-normalization algorithm, like the one presented in [11],
in order to render the result non-overlapping. Since this implies adding a sequential step at the end of the algorithm, slowing it’s performance, we recommend using this algorithm only if computations are known not to be error-prone.

5 Warp-synchronous GPU implementation

GPUs are highly multi-threaded SIMD architectures [12]. They are programmed in languages like CUDA and OpenCL, that expose a pure multi-thread programming model. Programmers describe compute kernels as a single program run by many fine-grained threads. The compiler and hardware group these threads into so-called warps, containing 32 threads on current Nvidia architectures. Threads inside a warp run in lockstep and share a single control flow, and their instructions are executed on SIMD units, with one thread per lane.

This implicit SIMD model is equivalent to explicit SIMD: a GPU program can be also understood as computations on vectors from the point of view of a warp. This enables direct implementation of the data-parallel algorithms we propose. Recent additions to the available hardware primitives make this warp-synchronous programming style particularly efficient [10].

Warp vote instructions perform boolean reductions across all threads within a warp. For instance, they can check whether a condition holds for all the threads, or for any of the threads of the warp. We use it to implement the loop exit conditions of the multiplication Algorithm 2.

Warp shuffle instructions allow arbitrary communication between threads in a warp, without having to go through memory. They are analogous to shuffle or permute instruction in explicit SIMD instruction sets [9]. We use them to shift vector components to propagate the errors across expansion terms, and to insert and extract scalar values inside vectors.

Our implementation targets GPUs with compute capability 3.0 or above, that offer the shuffle instruction. The code was written using CUDA C, using double-precision numbers (i.e. \( p = 53 \)). We illustrate the warp-synchronous implementation of Algorithm 2 for FP expansion multiplication in Fig. 5. The code appears from a single thread’s perspective, but it runs in parallel and it takes decisions based on the vector lane within an expansion (i.e. threadIdx.x). We use the shuffle instructions, that we extended for use with the double type:

- \( \text{shfl}_\text{up}(x, n, R) \) and \( \text{shfl}_\text{down} \) shift components respectively upward or downward by \( n \) positions within each \( R \)-element vector;
- \( \text{shfl} \) reads an arbitrary vector component within each vector lane.

Although we present here a version of the code that is parameterized by only one parameter, \( R \), in our actual implementation we use different templates for inputs

```c
template<int R>
__device__ double parallelMul(double x, double y){
    int lane = threadIdx.x; // Index within expansion
double s = 0., r = 0., y_i, p, s, e, ep;
for(int i=0; i<R-1; i++) {
    y_i = shfl(y, i, R); // Broadcast y_i
    p = TwoProd(x, y_i, &e);
    s = TwoSum(s, p, &ep);
    double tmp = shfl(s, 0, R);
    if(lane == i) r = tmp; // Save s_0 to r_i
    s = shfl_down(s, 1); // Shift left
    if(lane == R-1) s = 0.;
}
while(__any(e != 0.)) { // Accumulate e
    s = TwoSum(s, e, &e);
    e = shfl_up(e, 1, R); // Shift right
    if(lane == 0) e = 0.;
}
while(__any(ep != 0.)) { // Accumulate ep
    s = TwoSum(s, ep, &ep);
    ep = shfl_up(ep, 1, R); // Shift right
    if(lane == 0) ep = 0.;
}
}s = TwoSum(s, e, &e);
ep = shfl_up(ep, 1, R); // Shift right
if(lane == 0) ep = 0.;
}and output, meaning that we allow static generation of any input-output precision combinations.

We exploit both the parallelism that exist between expansion terms and across different expansions. To benefit from the SIMD execution and intra-warp communication primitives, all terms in a given expansion have to be computed by threads of the same warp. As warps have 32 threads on Nvidia architectures, the maximal supported expansion size is 32. Smaller expansions are packed together inside warps. Although this approach works with any expansion size \( R \) between 1 and 32 using appropriate padding, we recommend using power of two sizes, which allow filling the whole warp.

6 Comparison and discussion

In this section we present performance measurements obtained on a Tesla K40 GPU, using the CUDA 7.5 software architecture. We measure throughput on embarrassingly-parallel computations. The values are given in million of operations per second (Mop/s). By one op we understand one operation using extended precision. The tests were done using random generated examples, running on 1024 blocks each with 1024, 512 or 256 execution threads, depending on the expansion size and the required resources to run the algorithms.

To analyze the effect of parallelism on memory footprint, we consider two different shared memory usage scenarios: one best case that assumes the application uses no intermediate data outside of the registers used for the computation, and one worst case where the application
uses 256 bytes of CUDA shared memory for each term of the expansion.

For comparison, we report to Bailey’s GQD library [8] and CAMPARY [11], which offer extended precision using FP expansions on GPU. The arithmetic algorithms they employ are not by themselves parallel. Moreover, the GQD library is limited to double-double and quad-double precisions, and the algorithms used in the implementation are not provided with any correctness proof and there is no specific error bound given. CAMPARY is a recent software, that comes with correctness proofs and results guaranteed within a certain error bound.

In Table 1 we show the addition algorithm’s performance and in 2 the multiplication’s one for the best-case, no-memory configuration.

<table>
<thead>
<tr>
<th>$R$</th>
<th>Safe (Alg. 1)</th>
<th>Fast</th>
<th>CAMPARY</th>
<th>QD</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>6941</td>
<td>11914</td>
<td>3418</td>
<td>41549</td>
</tr>
<tr>
<td>4</td>
<td>1114</td>
<td>4667</td>
<td>1448</td>
<td>5310</td>
</tr>
<tr>
<td>8</td>
<td>314.4</td>
<td>2244</td>
<td>493.6</td>
<td>*</td>
</tr>
<tr>
<td>16</td>
<td>73.89</td>
<td>830.3</td>
<td>138.6</td>
<td>*</td>
</tr>
<tr>
<td>32</td>
<td>10.57</td>
<td>191.4</td>
<td>41.77</td>
<td>*</td>
</tr>
</tbody>
</table>

Even in this worst-case embarrassingly-parallel setup, the performance of data-parallel addition algorithm is competitive with the best known sequential algorithms for smaller expansions: the parallelism comes at little cost in number of operations per expansion. Addition on larger expansions and multiplications suffer from parallelization overhead.

The benefits of exploiting the parallelism available within each expansion are fully realized when parallelism is constrained by internal memory usage. Tables 3 and 4 respectively present addition and multiplication results in the memory-constrained configuration.

<table>
<thead>
<tr>
<th>$R$</th>
<th>Algorithm 2</th>
<th>CAMPARY</th>
<th>QD</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>6484</td>
<td>11100</td>
<td>27390</td>
</tr>
<tr>
<td>4</td>
<td>756.2</td>
<td>1363</td>
<td>2726</td>
</tr>
<tr>
<td>8</td>
<td>135.2</td>
<td>47.19</td>
<td>*</td>
</tr>
<tr>
<td>16</td>
<td>25.07</td>
<td>12.10</td>
<td>*</td>
</tr>
<tr>
<td>32</td>
<td>2.63</td>
<td>2.62</td>
<td>*</td>
</tr>
</tbody>
</table>

The performance of data-parallel algorithms remains stable in this setup, while the performance of sequential algorithm decreases sharply with memory usage. Although the QD library remains faster on expansions of size 2 (double-double), the data-parallel algorithms significantly outperform their sequential counterparts for all larger expansions. The performance gap increases with the expansion size, eventually reaching an order of magnitude for 32-term expansions.

### 7 Conclusion

We presented data-parallel algorithms for addition and multiplication of floating-point expansions. By taking advantage of data parallelism within FP expansions, they are suitable for SIMD architectures. We present fast addition and multiplication algorithms, as well as a safe addition algorithm with rigorous error bounds. A GPU implementation of the algorithms using warp-synchronous programming in CUDA is already competitive with state-of-the-art sequential algorithm in an idealistic embarrassingly parallel setup. However, data-parallel algorithms really shine in the more realistic case of applications that manipulate a sizable amount of intermediate data, significantly outperforming sequential algorithms for expansions of size 4 and greater.

### References


### Table 4: Performance in Mop/s for FP expansion multiplication algorithms in the memory-constrained case with 256B shared memory per expansion term

<table>
<thead>
<tr>
<th>$R$</th>
<th>Algorithm 2</th>
<th>CAMPARY</th>
<th>QD</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>882.87</td>
<td>2093</td>
<td>3501</td>
</tr>
<tr>
<td>4</td>
<td>99.37</td>
<td>180.8</td>
<td>256.9</td>
</tr>
<tr>
<td>8</td>
<td>18.05</td>
<td>7.06</td>
<td>*</td>
</tr>
<tr>
<td>16</td>
<td>4.04</td>
<td>0.84</td>
<td>*</td>
</tr>
<tr>
<td>32</td>
<td>0.23</td>
<td>0.096</td>
<td>*</td>
</tr>
</tbody>
</table>