

# Multiple Precision Arithmetic Versions of SDP solvers; SDPA-GMP, SDPA-QD and SDPA-DD

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## Collaborators

The SDPA project members in alphabetic order with **WAKI, Hayato**

- **FUJISAWA, Katsuki**
- FUKUDA, Mituhiro
- FUTAKATA, Yoshiaki
- KOBAYASHI, Kazuhiro
- KOJIMA, Masakazu
- NAKATA, Kazuhide
- (NAKATA, Maho)
- YAMASHITA, Makoto



# Outline

- 1 Introduction
  - Abstract
  - What is number?
  - Semidefinite programming
  - Necessity of accurate solver
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[Nakata-Nakatsuji-Ehara-Fukuda-Nakata-Fujisawa, J. Chem. Phys. 114, 8282 (2001)]
- Such problems require very high accuracy to SDP; relative gap  $< 1.0 \times 10^{-8}$
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Do not think seriously. Take it easy!

Keep it sweet and simple

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# What is number?

There are several kinds of numbers.

- Natural number:  $0, 1, 2, 3, 4, \dots$
- Integer:  $\dots, -3, -2, -1, 0, 1, 2, 3, 4, \dots$
- Rational number:  $a/b$ , where  $a, b$  are relatively prime
- Real number: convergence of Cauchy series.  
 $\{x_n : x_n \in \mathbb{Q}\}_{n=0,1,\dots}$  s.t.  $\forall \epsilon > 0, \exists N, \forall n, m > N \rightarrow |x_n - x_m| < \epsilon$   
defines a real number  $x$ .
- Complex number:  $z = a + bi$ : two real numbers with  $i$ .
- **floating point number**: designed for computers, subset of rational numbers.



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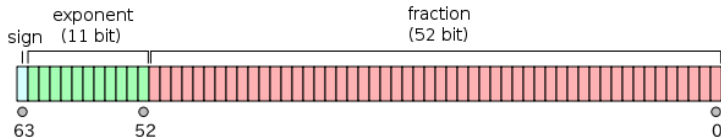
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# IEEE 754: Standard for Binary Floating-Point Arithmetic

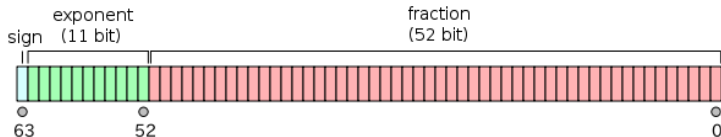
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- Very well designed we feel as if we treat real numbers.
- IEEE 754 double is expressed in 64-bit (= 8 bytes)



- $a = \pm \left( \frac{1}{2} + \frac{d_2}{2^2} + \frac{d_3}{2^3} + \dots + \frac{d_{52}}{2^{52}} \right) \times 2^e$ ,  $d = 0$  or  $1$ ,  
 $e = -1022 \sim 1023$
- about 16 significant digits ( $\log_{10} 2^{53} = 15.955$ ).
- Implemented for popular CPUs.

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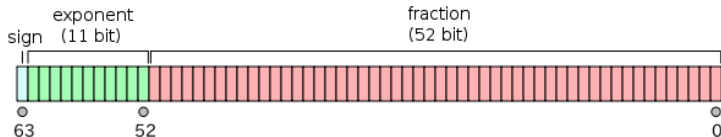
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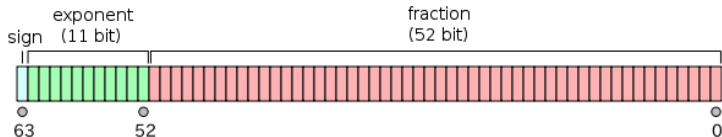


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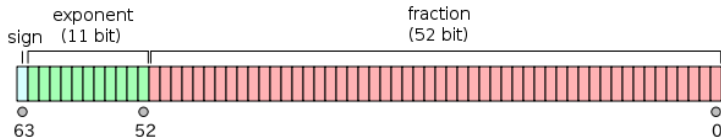
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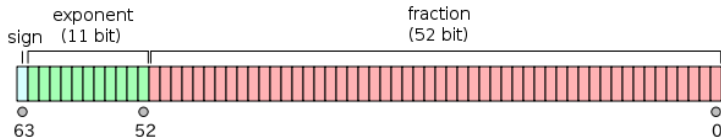
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# Semidefinite programming

$$\left\{ \begin{array}{ll}
 \text{primal} & \text{minimize:} & A_0 \bullet X \\
 & \text{s.t.:} & A_i \bullet X = b_i \quad (i = 1, 2, \dots, m) \\
 & & X \geq 0 \\
 \text{dual} & \text{maximize:} & \sum_{i=1}^m b_i z_i \\
 & \text{s.t.:} & \sum_{i=1}^m A_i z_i + Y = A_0 \\
 & & Y \geq 0
 \end{array} \right.$$

$A_i$  is  $n \times n$  real symmetric matrices,  $X$   $n \times n$  real symmetric variable matrix,  $b_i$  are constant vectors of  $m$ -dimension,  $Y$  is  $n \times n$  a real symmetric variable matrix,  $X \bullet Y := \sum X_{ij} Y_{ij}$ .  $X \geq 0$  means  $X$  is positive semidefinite.

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# Do we need to solve SDP problems *accurately*?

There are some questions for SDP results.

- Some SDPs are hard to solve. The results may have large gaps, not feasible.
- Simply we may not trust the results: “Strange Behaviors of Interior-point Methods for Solving Semidefinite Programming Problems in Polynomial Optimization” [Waki-Nakata-Muramatsu submitted]
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- IEEE 754 double arithmetic: done in 16 significant digits.  
accuracy losses in manipulations

$$1 \oplus 1.0 \times 10^{-17} = 1$$

- Condition number of matrix  $A$ ;  $\|A\| \|A^{-1}\|$ . when it becomes  $10^{16}$ , solution to the linear equation is inaccurate with IEEE 754 double.
- $X \bullet Y = 0$  at the optimum (complementarity slackness theorem for SDP)  
variable matrix becomes singular at the optimum; condition number becomes infinite!



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- $Z^{-1}$ : Primal-dual interior point method calculates  $Z^{-1}$ ; *“It is seldom necessary to compute the inverses of matrix explicitly, and it is certainly not recommended as a means of solving linear systems.”* by LAPACK Users’ Guide Third Edition, p.14.
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# A brute force method for accurate SDP solutions

- Use multiple precision arithmetic; GMP, QD rather than IEEE 754 double.
- Simple answer to obtain high accuracy.
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# What is GMP?

- GMP is a free library for arbitrary precision arithmetic, operating on signed integers, rational numbers, and **floating point numbers**.
- significant digits: arbitrary (I usually use 60 ~ 72 digits)



## Strategy and features

- Using existing multiple precision libraries.
- Based on SDPA; <http://sdpa.indsys.chuo-u.ac.jp/sdpa/>
- No changes in algorithm.
- Changes from SDPA should be minimal to reduce the maintenance cost.
- Matrix-vector manipulations and eigenvalues etc. → **Multiple precision version of LAPACK and BLAS.**
  - 49 routines from MPACK; Rpotrf (dpotrf.f; cholesky), Rsyev (dsyev.f eigenvalue), Rsterf, Rsteqr (dsterf.f, dsteqr.f) etc..
- Introduction of “**precision**” parameter; controls number of significant bits used in the calculations.
- Actually I did was **replacing “double” to “mpf\_class”** carefully.

## Another MP library: Quad-Double library

- Usually quadruple or octuple precision are enough.
- Double-Double and Quad-Double Arithmetic; by Y. Hida, Xiaoye S. Li, David H Bailey, and **faster** than GMP.
- Four/two unevaluated IEEE 754 double  $\sim$  approx octuple/quadruple precision.

$$A = (a_0, a_1, a_2, a_3)$$

- Utilize exact transformations [Dekker, Knuth, Priest, Shewcheck].

$$a = x \oplus y, b = x + y - (x \oplus y)$$

Error by IEEE754 add  $x \oplus y$  can be *exactly* evaluated.

- Replace “mpf\_class” to “dd\_real” and “qd\_real”  $\rightarrow$  SDPA-QD, SDPA-DD.

## What is MPACK?

- **MPACK is a multiple precision version of BLAS and LAPACK. <http://mplapack.sourceforge.net/>**
- What is the BLAS? The BLAS (Basic Linear Algebra Subprograms) are routines that provide standard building blocks for performing basic vector and matrix operations.
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## What is MPACK?

- MPACK is a multiple precision version of BLAS and LAPACK. <http://mplapack.sourceforge.net/>
- What is the BLAS? The BLAS (Basic Linear Algebra Subprograms) are routines that provide standard building blocks for performing basic vector and matrix operations.
- What is LAPACK? This provides routines for solving systems of simultaneous linear equations, least-squares solutions of linear systems of equations, eigenvalue problems, and singular value problems.
- Written C++, but look very similar to reference BLAS implementation.
- Very portable and no optimization at this moment.
- Pretty good compatibility with BLAS and LAPACK.

# MBLAS Rgemm.cpp and BLAS dgemm.f

## Rgemm.cpp

```
//Start the operations.
if (notb) {
  if (nota) {
    //Form C := alpha*A*B + beta*C.
    for (int j = 0; j < n; j++) {
      if (beta == Zero) {
        for (int i = 0; i < m; i++) {
          C[i + j * ldc] = Zero;
        }
      } else if (beta != One) {
        for (int i = 0; i < m; i++) {
          C[i + j * ldc] = beta * C[i + j * ldc];
        }
      }
      for (int l = 0; l < k; l++) {
        if (B[l + j * ldb] != Zero) {
          temp = alpha * B[l + j * ldb];
          for (int i = 0; i < m; i++) {
            C[i + j * ldc] =
              C[i + j * ldc] + temp * A[i + l * lda];
          }
        }
      }
    }
  } else {
    //Form C := alpha*A'*B + beta*C.

```

## dgemm.f

```
*
* Start the operations.
*
IF (NOTB) THEN
  IF (NOTA) THEN
*
* Form C := alpha*A*B + beta*C.
*
    DO 90 J = 1,N
      IF (BETA.EQ.ZERO) THEN
        DO 50 I = 1,M
          C(I,J) = ZERO
        CONTINUE
      ELSE IF (BETA.NE.ONE) THEN
        DO 60 I = 1,M
          C(I,J) = BETA*C(I,J)
        CONTINUE
      END IF
      DO 80 L = 1,K
        IF (B(L,J).NE.ZERO) THEN
          TEMP = ALPHA*B(L,J)
          DO 70 I = 1,M
            C(I,J) = C(I,J) + TEMP*A(I,L)
          CONTINUE
        END IF
      CONTINUE
    CONTINUE
  ELSE
*
* Form C := alpha*A'*B + beta*C
*

```

## How MBLAS is used in SDPA-GMP?

From `sdpa_linear.cpp` from SDPA-GMP 7.1.2.

```
if (scalar==NULL) {
    scalar = &MONE;
    // scalar is local variable
}
// The Point is the first argument is "Transpose".
Rgemm("Transpose", "NoTranspose", retMat.nRow, retMat.nCol, aMat.nCol,
      *scalar, aMat.de_ele, aMat.nCol, bMat.de_ele, bMat.nRow,
      0.0, retMat.de_ele, retMat.nRow);
break;
case DenseMatrix::COMPLETION:
    rError("no support for COMPLETION");
    break;
}

return _SUCCESS;
```

# Results (I)

Some results from SDPLIB, on Opteron 250 (2.4GHz), 16G Mem, FreeBSD 7/amd64.  
 "precision" is 250 for GMP.

instance	arch8(GMP)	arch8(QD)	arch8(DD)	arch8(double)
iter	47	47	37	25
rel. gap	3.57e - 31	3.58e - 31	3.80e - 21	1.65e - 08
p feas. error	3.11e - 76	1.02e - 61	5.05e - 29	1.14e - 12
d feas. error	5.66e - 72	9.01e - 52	4.85e - 21	1.10e - 07
time (s)	634.766	497.289	55.445	9.35

instance	mcp500-4(GMP)	mcp500-4(QD)	mcp500-4(DD)	mcp500-4(double)
iter	38	38	28	15
rel. gap	1.36e - 31	1.36e - 31	1.36e - 21	1.16e - 08
p feas. error	1.28e - 76	6.08e - 64	6.41e - 31	4.88e - 15
d feas. error	1.67e - 75	7.72e - 59	1.68e - 28	1.02e - 13
time (s)	5711.6	4678.1	455.0	10.2

instance	maxG32(GMP)	maxG32(QD)	maxG32(DD)	maxG32(double)
iter	40	40	30	17
rel. gap	2.07e - 31	2.07e - 31	2.04e - 21	1.65e - 08
p feas. error	1.74e - 76	1.09e - 64	1.23e - 31	1.14e - 12
d feas. error	1.90e - 72	2.47e - 53	8.53e - 25	1.10e - 07
time (s)	348564.8	315969.5	30472.0	9.35





## Results (II) 1D-Hubbard model

1D Hubbard model Strong correlation limit:  $|U/t| \rightarrow \infty$ : [Nakata et al. JCP 2008]; with SDPA-GMP 6.

### Ground state energy of 1D Hubbard model

PBC, # of sites:4, # of electrons: 4, spin 0

U/t	SDPA (16 digits)	SDPA-GMP (60 digits)	fullCI
10000.0	0	$-1.1999998800000251 \times 10^{-3}$	$-1.199999880 \times 10^{-3}$
1000.0	$-1.2 \times 10^{-2}$	$-1.1999880002507934 \times 10^{-2}$	$-1.1999880002 \times 10^{-2}$
100.0	$-1.1991 \times 10^{-1}$	$-1.1988025013717993 \times 10^{-1}$	$-1.19880248946 \times 10^{-1}$
10.0	-1.1000	-1.0999400441222934	-1.099877772750
1.0	-3.3417	-3.3416748070259956	-3.340847617248

PBC, # of sites:6, # of electrons: 6, spin 0

U/t	SDPA (16 digits)	SDPA-GMP (60 digits)	fullCI
10000.0	0	$-1.7249951195749525 \times 10^{-3}$	$-1.721110121 \times 10^{-3}$
1000.0	$-1 \times 10^{-2}$	$-1.7255360310431304 \times 10^{-2}$	$-1.7211034713 \times 10^{-2}$
100.0	$-1.730 \times 10^{-1}$	$-1.7302157140594339 \times 10^{-1}$	$-1.72043338097 \times 10^{-1}$
10.0	-1.6954	-1.6953843276854447	-1.664362733287
1.0	-6.6012	-6.6012042217806286	-6.601158293375

## Results (III) Kissing number

Kissing number from A New Library of Structured Semidefinite Programming Instances; the optimal values *were* uncertain or only known with low accuracy. Powered by **Fujisawa-san** (2008/12/21); precision is 128bit for GMP.

instance	opt (double)	opt (GMP)
kissing_3_10_10	-11.4385	-11.43814328
kissing_4_10_10	-23.14	-23.13553364
kissing_5_10_10	-44.15	-44.158868754
kissing_6_10_10	-77.9	-77.912852357
kissing_7_10_10	-134.3	-134.32853967
kissing_8_10_10	-238.929	-238.99981527
kissing_9_10_10	-365	-365.21946909
kissing_10_10_10	-562.9	-562.89594739
kissing_11_10_10	-889.74	-889.74203646
kissing_12_10_10	-1369.485	-1369.5287720

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# Summary

- We developed multiple precision version of SDP solver. SDPA-GMP, SDPA-QD and SDPA-DD.
- Can solve SDPs very accurately.
- MPACK 0.0.9: Multiple precision version of LAPACK/BLAS: development ongoing.
- Outlook
  - Faster SDPA-GMP, QD, DD and MPACK, parallel and multicore versions.
  - More routines for MPACK.
  - Higher accuracy to SDPA; minimal use of multiple precision arithmetic.

