

# User Guide of CKTSO

(version 20231101)

Xiaoming Chen ([chenxiaoming@ict.ac.cn](mailto:chenxiaoming@ict.ac.cn))

## Contents

1. Introduction .....	2
2. License Key .....	2
3. System Requirements .....	3
4. Matrix Format.....	3
5. C/C++ Functions .....	4
5.1 Create Solver .....	4
5.2 Destroy Solver .....	5
5.3 Matrix Analysis.....	6
5.4 Factorize with Pivoting.....	7
5.5 Refactorize without Pivoting.....	9
5.6 Solve.....	10
5.7 Solve Multiple Vectors.....	11
5.8 Sort Factors .....	13
5.9 Calculate Workloads.....	14
5.10 Clean Up Garbage.....	15
5.11 Calculate Determinant .....	16
5.12 Factorize and Solve .....	16
5.13 Refactorize and Solve.....	17
6. Parameters .....	18
6.1 Input Parameters .....	18
6.2 Output Parameters .....	20
7. Using CKTSO Libraries.....	21
8. Using CKTSO in Circuit Simulators .....	21

## 1. Introduction

CKTSO is a high-performance parallel sparse direct solver specially designed for **SPICE-based circuit simulation**. It solves  $Ax = b$  where  $A$  is square and highly sparse. The software is written in pure C and provides both C and C++ interfaces. Dynamic-link libraries for Windows and Linux are provided.

CKTSO is the successor of our previous work, NICSLU (<https://github.com/chenxm1986/nicslu>). CKTSO uses many similar techniques to NICSLU. However, CKTSO integrates some novel techniques and shows higher performance, better scalability and less memory usage than NICSLU, while NICSLU provides more functionalities. The most important features of CKTSO include

- a) a new pivoting-reduction technique that significantly improves the performance and scalability of LU factorization with pivoting;
- b) a new memory allocation strategy that reduces memory usage;
- c) parallel forward and backward substitutions;
- d) a novel nested dissection ordering method, which reduces about 10X floating-point operations for post-layout/mesh-style circuits, and it also produces fewer floating-point operations than METIS;
- e) several novel minimum degree ordering variants, which reduce about 30-40% floating-point operations compared with the approximate minimum degree;
- f) an adaptive numerical kernel selection method.

CKTSO solves a sparse linear system through three main steps: **symbolic analysis, factorization, and solving**. Symbolic analysis tries to order the matrix to minimize fill-ins that will be generated in factorization. In circuit simulation, symbolic analysis is usually executed only once. The factorization step factorizes the matrix into the product of a lower triangular matrix and an upper triangular matrix, i.e.,  $A = LU$ . Partial pivoting can be performed to ensure the numerical stability. The solving step finds the solution of the linear system through two triangular system solvers. i.e.,  $Ly = b$  and  $Ux = y$ .

Some algorithms of CKTSO are described in the following paper.

- Xiaoming Chen, "Numerically-Stable and Highly-Scalable Parallel LU Factorization for Circuit Simulation", in 2022 International Conference On Computer Aided Design (ICCAD'22).

## 2. License Key

Running CKTSO requires a valid license key file. The license key file must be named "cktso.lic" and put **together with the library file** (\*.dll on Windows and \*.so on Linux). It is a pure text file. Do not edit any content of the license key file, or it may be invalidated.

### 3. System Requirements

Table 1 lists the basic software and hardware requirements. CKTSO libraries can be run on Windows or Linux. For Linux, libraries for CentOS and Ubuntu are provided, and they are compatible with many mainstream Linux distributions. CKTSO libraries are provided as x64 dynamic-link libraries which can run on x64 CPUs. Pure x86 libraries are not provided. CKTSO uses AVX2 and FMA instructions. If the CPU does not support such instructions, an error will occur when running CKTSO libraries.

Table 1. Basic software and hardware requirements

Operating system	Windows 7 or higher, or mainstream Linux distributions
CPU	x86_64, with AVX2 and FMA instructions supported

### 4. Matrix Format

The input format of CKTSO is the **compressed sparse row (CSR)** format. CSR uses an integer  $n$  and three arrays  $ap[]$ ,  $ai[]$  and  $ax[]$  to represent a sparse matrix.  $n$  is the matrix dimension. Array  $ap[]$  of length  $n+1$  stores the row pointers. Specifically,  $ap[i]$  stores the start position of every row in  $ai[]$  and  $ax[]$ . Array  $ai[]$  of length  $ap[n]$  stores the column indexes. Array  $ax[]$  of length  $ap[n]$  stores the matrix values (one-to-one corresponding to the elements of array  $ai[]$ ). Figure 1 illustrates an example of CSR, in which the matrix is of dimension 4.

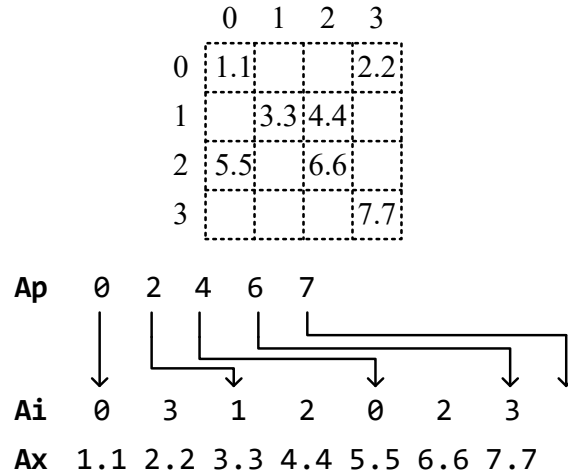


Figure 1: Example of CSR.

CKTSO does not need the column indexes in each row to be sorted. From version 20231101, duplicated entries in CSR are allowed. However, the existence of duplicated entries will introduce additional latency and memory overheads. From version 20221214, **CKTSO natively supports both real-**

**number and complex-number matrices.** The performance of complex-number computation has not been fully optimized.

## 5. C/C++ Functions

CKTSO provides both C and C++ interfaces. Both 32-bit and 64-bit integer versions are provided, and the latter involves a "\_L" flag in the function name or object name. The 64-bit interface uses 64-bit integers to represent the indexes of the input matrix. The internal data structures always use 64-bit integers to store the LU factors, even for the 32-bit interface.

Both the C and C++ interfaces involve an ICktSo (struct \_\_cktso\_dummy \*) or ICktSo\_L (struct \_\_cktso\_l\_dummy \*) object. An object instance is created by CKTSO\_CreateSolver, CKTSO\_L\_CreateSolver, CKTSO\_CreateSolverNoCheck, or CKTSO\_L\_CreateSolverNoCheck and passed as the first argument to other C functions. Users may call its member functions in the C++ environment. The member functions of the object have one-to-one correspondence with the C functions, e.g., \_\_cktso\_dummy::Analyze corresponds to CKTSO\_Analyze.

All CKTSO functions (including both C and C++ functions) return an integer to indicate the error. Zero means no error and negative values indicate errors. The meaning of the return code is listed in Table 2.

Table 2: Return code.

0	Successful	-1	Invalid instance handle
-2	Argument error	-3	Invalid matrix data
-4	Out of memory	-5	Structurally singular
-6	Numerically singular	-7	Threads-related error
-8	Matrix has not been analyzed	-9	Matrix has not been factorized
-10	Function is not supported	-11	File cannot be open
-12	Integer overflow	-13	Resource leak
-99	License error	-100	Unknown error

### 5.1 Create Solver

```
int CKTSO_CreateSolver
(
    ICktSo *inst,
    int **iparm,
    const long long **oparm
);
```

```
int CKTSO_L_CreateSolver
(
    ICKtSo_L *inst,
    int **iparm,
    const long long **oparm
);
```

```
int CKTSO_CreateSolverNoCheck
(
    ICKtSo *inst,
    int **iparm,
    const long long **oparm
);
```

```
int CKTSO_L_CreateSolverNoCheck
(
    ICKtSo_L *inst,
    int **iparm,
    const long long **oparm
);
```

This function creates the solver instance. The created instance can be used in both C and C++ environments.

- The parameter `inst` returns the pointer to the created instance.
- The parameters `iparm` and `oparm` return the pointers to the internal input parameter and output parameter arrays, respectively. They can be NULL if not needed. If so, there will be no chance to change the configurations or retrieve the statistical information of the solver. Do not free `iparm` and `oparm`. The arrays will be freed when the solver instance is destroyed.

`CKTSO_CreateSolver` or `CKTSO_L_CreateSolver` checks whether the CPU supports AVX2 and FMA instructions, and returns -10 if these instructions are not supported. `CKTSO_CreateSolverNoCheck` or `CKTSO_L_CreateSolverNoCheck` does not perform such check. This is useful on virtual machines which do not support these instructions but the CPU actually supports them.

## 5.2 Destroy Solver

```
int CKTSO_DestroySolver
(
    ICKtSo inst
);
```

```
int CKTSO_L_DestroySolver
(
    ICKtSo_L inst
);
```

```
virtual int __cktso_dummy::DestroySolver
(
) = 0;
```

```
virtual int __cktso_l_dummy::DestroySolver
(
) = 0;
```

This function frees any data associated with the specified instance and also destroys the instance. Previously created threads will also exit. After that, the instance is invalid. Any created instance should be destroyed when it will no longer be used. Do not destroy an instance more than once.

### 5.3 Matrix Analysis

```
int CKTSO_Analyze
(
    ICKtSo inst,
    bool is_complex,
    int n,
    const int ap[],
    const int ai[],
    const double ax[],
    int threads
);
```

```
int CKTSO_L_Analyze
(
    ICKtSo_L inst,
    bool is_complex,
    long long n,
    const long long ap[],
    const long long ai[],
    const double ax[],
    int threads
);
```

```
virtual int __cktso_dummy::Analyze
(
    bool is_complex,
    int n,
    const int ap[],
    const int ai[],
    const double ax[],
    int threads
) = 0;
```

```

virtual int __cktso_1_dummy::Analyze
(
    bool is_complex,
    long long n,
    const long long ap[],
    const long long ai[],
    const double ax[],
    int threads
) = 0;

```

This function creates internal data for the matrix, reorders the matrix to minimize fill-ins, performs static symbolic factorization and also creates threads. It must be called before any factorization is called. In circuit simulation, it usually needs only once because the symbolic structure of the matrix is fixed during iterations.

- The parameter `is_complex` specifies whether the matrix is complex. For the complex case, all double-valued arrays should store complex numbers in an interleaved real-imaginary form. The complex-number data type can be defined as `double [2]`.
- The parameter `n` specifies the matrix dimension. Specifying a positive `n` will first destroy the previous matrix and perform analysis for the new matrix. Specifying `n=0` will only destroy the previous matrix, without creating the new matrix and doing analysis for the new matrix (the solver instance is still valid).
- The parameters `ap`, `ai` and `ax` specify the CSR arrays of the matrix. `ax` can be NULL if the values are unavailable when the matrix is analyzed.
- The parameter `threads` specifies the number of threads. The created threads will be used for matrix analysis, factorization, refactorization, sorting factors, and solving. They are managed as a thread pool and will not exit until the solver instance is destroyed or the matrix is destroyed. Specifying `threads=0` will use all physical CPU cores, and specifying `threads=-1` means using all logical CPU cores. The specified number of threads cannot exceed the number of logical CPU cores. The maximum number of threads is suggested to be the number of physical CPU cores (specifying `threads=0`). It is not suggested to use more threads than the number of physical cores. Depending on the hardware and operating system, CKTSO may not retrieve the correct number of physical cores, so it is highly recommended that the number of threads is explicitly specified.

## 5.4 Factorize with Pivoting

```

int CKTSO_Factorize
(
    ICKtSo inst,
    const double ax[],
    bool fast
);

int CKTSO_L_Factorize
(
    ICKtSo_L inst,
    const double ax[],
    bool fast
);

virtual int __cktso_dummy::Factorize
(
    const double ax[],
    bool fast
) = 0;

virtual int __cktso_l_dummy::Factorize
(
    const double ax[],
    bool fast
) = 0;

```

This function factorizes the reordered matrix into LU factors with partial pivoting. The factorization uses the threads created in matrix analysis. CKTSO has a thread control technique that can automatically judge whether using multiple threads is useful. If not, it will automatically use a single thread, even if multiple threads have been created. Except for the first factorization, CKTSO employs a fast pivoting-reduction technique which can significantly improve the performance of matrix factorization but the numerical stability is still maintained. CKTSO also utilizes a sparsity-oriented algorithm selection method to enhance the factorization performance for different sparsities.

- The parameter `ax` specifies the array storing the matrix values, which is of length `ap[n]` specified in matrix analysis.
- The parameter `fast` specifies whether fast factorization based on pivoting reduction is enabled.

Please note that **subsequent factorizations are generally much faster than the first-time factorization, if fast factorization is enabled**. In the best case, the performance and scalability of fast factorization are almost same as those of refactorization without pivoting. However, in some extreme cases, the fast factorization technique may lead to more fill-ins. It is suggested that **the first factorization of each independent simulation method (e.g., a DC simulation method like GMIN stepping or**



**pseudo-transient, the entire transient simulation, etc.) should disable fast factorization.** Please refer to Section 8 for more details.

## 5.5 Refactorize without Pivoting

```
int CKTSO_Refactorize
(
    ICktSo inst,
    const double ax[]
);

int CKTSO_L_Refactorize
(
    ICktSo_L inst,
    const double ax[]
);

virtual int __cktso_dummy::Refactorize
(
    const double ax[]
) = 0;

virtual int __cktso_l_dummy::Refactorize
(
    const double ax[]
) = 0;
```

This function refactorizes the matrix without pivoting. It should be called after factorization with pivoting has been called. It reuses the pivoting order and the symbolic structure of the LU factors obtained in the last factorization with pivoting. The refactorization uses the threads created in matrix analysis. CKTSO has a thread control technique that can automatically judges whether using multiple threads is useful. If not, it will automatically use a single thread, even if multiple threads have been created.

In circuit simulation, there are usually many Newton-Raphson iterations. When Newton-Raphson iterations are converging, the matrix values tend to change slowly. In this situation, factorizing the matrix without pivoting is generally numerically stable. This function provides an opportunity to improve the solver performance in circuit simulation by utilizing the features of Newton-Raphson iterations. To judge whether Newton-Raphson iterations are converging, one can simply check the difference between the solutions of two adjacent iterations. See Section 8 for details.

- The parameter `ax` specifies the array storing the matrix values, which is of length `ap[n]` specified in matrix analysis.

**Every time after the symbolic pattern of the LU factors has been changed (factorization with pivoting has been called), the first call of this function has an additional scheduler initialization step,**

which causes some additional time.

## 5.6 Solve

```
int CKTSO_Solve
(
    ICKtSo inst,
    const double b[],
    double x[],
    bool force_seq,
    bool row0_column1
);

int CKTSO_L_Solve
(
    ICKtSo_64 inst,
    const double b[],
    double x[],
    bool force_seq,
    bool row0_column1
);

virtual int __cktso_dummy::Solve
(
    const double b[],
    double x[],
    bool force_seq,
    bool row0_column1
) = 0;

virtual int __cktso_l_dummy::Solve
(
    const double b[],
    double x[],
    bool force_seq,
    bool row0_column1
) = 0;
```

This function performs forward and backward substitutions to solve the linear system, after the matrix has been factorized. CKTSO supports sequential or parallel solving. CKTSO has a thread control technique that can automatically judges the best number of threads for parallel solving.

- The parameter `b` specifies the right-hand-side vector.
- The parameter `x` specifies the solution vector. The address of array `x[ ]` may be same as the address of array `b[ ]`. If so, the solution vector is overwritten on the right-hand-side vector.
- The parameter `force_seq` specifies whether to force CKTSO to perform

sequential solving. However, even if forced sequential solving is not enabled, CKTSO does not necessarily perform parallel solving. Instead, CKTSO has a mechanism to determine whether to perform parallel solving and also the optimal number of threads automatically. Please refer to Section 8 for more details.

- The parameter `row0_column1` specifies whether the matrix is transposed (i.e., whether to solve  $A^T x = b$ ). Please note that CKTSO uses the **row-major CSR format** by default. This means that in the transposed mode the matrix should be stored in the column-major format.

**Every time after the symbolic pattern of the LU factors has been changed (factorization with pivoting has been called), the first call of this function has an additional scheduler initialization step, which causes about 2-5X time of the sequential solving time.**

Table 2: Performance of parallel solving.

	Windows	Linux
Row-major order	Generally most matrices have speedups but a few do not. Parallel solving is <b>recommended</b> in this scenario.	Almost all matrices have speedups. Parallel solving is <b>recommended</b> in this scenario.
Column-major order	Some matrices have speedups but some do not. Parallel solving is <b>not recommended</b> in this scenario.	Generally most matrices have speedups but a few do not. Parallel solving is <b>recommended</b> in this scenario.

The performance of parallel solving varies on different operating systems and for different matrix storage formats. Due to the much stronger dependency in parallel solving for matrices stored in the column-major format, column-mode parallel solving usually has lower performance than row-mode parallel solving. Table 2 summarizes a general description of the performance of parallel solving, as well as our recommendations.

## 5.7 Solve Multiple Vectors

```

int CKTSO_SolveMV
(
    ICKtSo inst,
    size_t nrhs,
    const double b[],
    size_t ld_b,
    double x[],
    size_t ld_x,
    bool row0_column1
);

```

```

int CKTSO_L_SolveMV
(
    ICKtSo_64 inst,
    size_t nrhs,
    const double b[],
    size_t ld_b,
    double x[],
    size_t ld_x,
    bool row0_column1
);

```

```

virtual int __cktso_dummy::SolveMV
(
    size_t nrhs,
    const double b[],
    size_t ld_b,
    double x[],
    size_t ld_x,
    bool row0_column1
) = 0;

```

```

virtual int __cktso_l_dummy::SolveMV
(
    size_t nrhs,
    const double b[],
    size_t ld_b,
    double x[],
    size_t ld_x,
    bool row0_column1
) = 0;

```

This function performs forward and backward substitutions for solving multiple vectors (i.e., solving  $\mathbf{Ax}_i = \mathbf{b}_i$ , where  $i = 1, 2, \dots, nrhs$ ), after the matrix has been factorized. If multiple threads have been created when analyzing the matrix, each thread will solve a vector independently; otherwise

CKTSO just solves all vectors sequentially.

- The parameter `nrhs` specifies the number of vectors to be solved.
- The array `b[ ]` is of length  $n \times \text{ld\_b}$ , and it stores the right-hand-vectors vector by vector on input. `ld_b` is the leading dimension of `b[ ]`. If `ld_b=0`, `ld_b=n`. In other cases, `ld_b` cannot be smaller than `n`.
- The array `x[ ]` is of length  $n \times \text{ld\_x}$ , and it stores the solution vectors vector by vector on output. `ld_x` is the leading dimension of `x[ ]`. If `ld_x=0`, `ld_x=n`. In other cases, `ld_x` cannot be smaller than `n`. The address of array `x[ ]` may be same as the address of array `b[ ]`. If so, the solution vectors are overwritten on the right-hand-side vectors.
- The parameter `row0_column1` specifies whether the matrix is transposed (i.e., whether to solve  $A^T x = b$ ).

The meaning of *leading dimension* is illustrated in Figure 2. When accessing a submatrix in a full matrix, using leading dimension is helpful. In short, leading dimension is the size of the first dimension of the full matrix, with respect to the storage order. If the matrix is stored in the row-major order, to get the next row head of the submatrix, one simply adds the leading dimension to the current row head. It is same for column-major stored matrices, if submatrices are accessed in the column order.

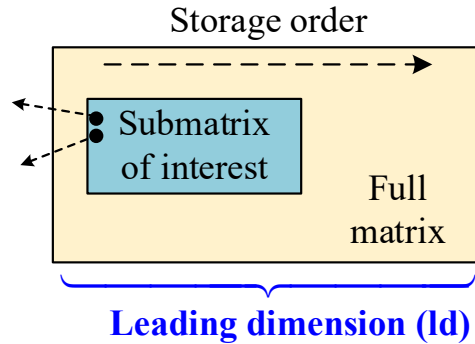


Figure 2: Leading dimension.

## 5.8 Sort Factors

```
int CKTSO_SortFactors
(
    ICKtSo inst,
    bool sort_values
);

int CKTSO_L_SortFactors
(
    ICKtSo_L inst,
    bool sort_values
);
```

```

virtual int __cktso_dummy::SortFactors
(
    bool sort_values
) = 0;
virtual int __cktso_l_dummy::SortFactors
(
    bool sort_values
) = 0;

```

This function sorts each row of the LU factors. It is typically used after factorization. Due to the factorization algorithm, the indexes of the LU factors are not guaranteed to be in order. With sorted factors, the cache efficiency of subsequent refactorization and solving processes may be improved, but the improvement is typically small on CPU. Sorting factors is a necessary step for the GPU acceleration module. It is strongly recommended to sort the factors before initializing the GPU acceleration module.

- The parameter `sort_values` specifies whether to sort values as well. If not, only indexes are sorted. If values are not sorted, a subsequent solving will return an error unless a factorization or refactorization has been called before solving.

## 5.9 Calculate Workloads

```

int CKTSO_Statistics
(
    ICKtSo inst,
    long long *factor_flops,
    long long *solve_flops,
    long long *factor_mem,
    long long *solve_mem,
    bool row0_column1
);
int CKTSO_L_Statistics
(
    ICKtSo_L inst,
    long long *factor_flops,
    long long *solve_flops,
    long long *factor_mem,
    long long *solve_mem,
    bool row0_column1
);

```

```
virtual int __cktso_dummy::Statistics
(
    long long *factor_flops,
    long long *solve_flops,
    long long *factor_mem,
    long long *solve_mem,
    bool row0_column1
) = 0;
```

```
virtual int __cktso_l_dummy::Statistics
(
    long long *factor_flops,
    long long *solve_flops,
    long long *factor_mem,
    long long *solve_mem,
    bool row0_column1
) = 0;
```

This function calculates the number of floating-operations and memory access volumes of factorization and solving. It should be called after factorization with partial pivoting has been called.

factor\_flops cannot be NULL. Each of the other three parameters, solve\_flops, factor\_mem and solve\_mem, can be NULL if not required. No initialization for the four parameters is required. The memory access volumes are reported in bytes. The parameter row0\_column1 specifies whether to solve  $A^T x = b$ .

## 5.10 Clean Up Garbage

```
int CKTSO_CleanUpGarbage
(
    ICktSo inst
);
```

```
int CKTSO_L_CleanUpGarbage
(
    ICktSo_L inst
);
```

```
virtual int __cktso_dummy::CleanUpGarbage
(
) = 0;
```

```
virtual int __cktso_l_dummy::CleanUpGarbage
(
) = 0;
```

This function cleans up redundant memory. If factorization with partial pivoting has been called many times, especially when Newton-Raphson

iterations diverge, the memory usage may be more than required. This function can clean up such unnecessary memory consumption.

### 5.11 Calculate Determinant

```
int CKTSO_Determinant
(
    ICktSo inst,
    double *mantissa,
    double *exponent
);

int CKTSO_L_Determinant
(
    ICktSo_L inst,
    double *mantissa,
    double *exponent
);

virtual int __cktso_dummy::Determinant
(
    double *mantissa,
    double *exponent
) = 0;

virtual int __cktso_l_dummy::Determinant
(
    double *mantissa,
    double *exponent
) = 0;
```

This function calculates the determinant of the matrix, after the matrix has been factorized. The determinant is expressed in the form of  $\text{mantissa} \times 10^{\text{exponent}}$ , where  $1 \leq |\text{mantissa}| < 10$ .

- If the matrix is complex, the parameter *mantissa* points to a complex number, while *exponent* still points to a real value.

### 5.12 Factorize and Solve

```
int CKTSO_FactorizeAndSolve
(
    ICktSo inst,
    const double ax[],
    const double b[],
    double x[],
    bool row0_column1
);
```



```

int CKTSO_L_FactorizeAndSolve
(
    ICktSo_L inst,
    const double ax[],
    const double b[],
    double x[],
    bool row0_column1
);

virtual int __cktso_dummy::FactorizeAndSolve
(
    const double ax[],
    const double b[],
    double x[],
    bool row0_column1
) = 0;

virtual int __cktso_l_dummy::FactorizeAndSolve
(
    const double ax[],
    const double b[],
    double x[],
    bool row0_column1
) = 0;

```

This function factorizes the matrix and then solves the linear system. Whether to use fast factorization and parallel solving is determined by a heuristic method according to past factorizations.

### 5.13 Refactorize and Solve

```

int CKTSO_RefactorizeAndSolve
(
    ICktSo inst,
    const double ax[],
    const double b[],
    double x[],
    bool row0_column1
);

```

```

int CKTSO_L_RefactorizeAndSolve
(
    ICKtSo_L inst,
    const double ax[],
    const double b[],
    double x[],
    bool row0_column1
);

virtual int __cktso_dummy::RefactorizeAndSolve
(
    const double ax[],
    const double b[],
    double x[],
    bool row0_column1
) = 0;

virtual int __cktso_l_dummy::RefactorizeAndSolve
(
    const double ax[],
    const double b[],
    double x[],
    bool row0_column1
) = 0;

```

This function refactorizes the matrix and then solves the linear system. Whether to use parallel solving is determined by a heuristic method according to past factorizations.

## 6. Parameters

CKTSO maintains an input parameter array for behavior configurations and an output parameter array for information statistics. Input parameters are initialized to default values in CKTSO\_CreateSolver or CKTSO\_L\_CreateSolver. When calling CKTSO\_CreateSolver or CKTSO\_L\_CreateSolver, one has a chance to retrieve the pointers to the two internal arrays, which are in `**iparm` and `const long long **oparm`. By setting the input parameter array, the behavior of CKTSO may be configured. Most input parameters are effective if they are set before matrix analysis.

### 6.1 Input Parameters

`iparm[0]`: timer control. Zero means no timer. A positive value means using a high-precision timer while a negative value means using a low-precision timer. High-precision timer also involves higher overhead than low-precision timer. Only when `iparm[0]` is

- nonzero, `oparm[0]` to `oparm[3]` are effective. Default is zero (timer disabled).
- `iparm[1]`: pivoting tolerance. It is in millionth ( $1/1000000$ ). Default is 1000 (0.001 actually).
- `iparm[2]`: ordering method. CKTSO has 10 different ordering methods including 8 minimum degree variants and 2 nested dissection variants. Zero means selecting the best from the 10 methods. 1 to 10 mean using the corresponding single ordering method, where 1 to 8 correspond to minimum degree variants and 9 to 10 correspond to nested dissection variants. 11 means selecting the best from the 2 nested dissection methods. 12 to 18 mean selecting the best from `iparm[2]-10` minimum degree variants. For example, specifying `iparm[2]=15` means selecting the best from the minimum degree ordering methods 1 to 5. A negative value means no ordering (using the natural order). Default is zero (selecting best from all 10 methods).
- `iparm[3]`: threshold for dense node detection. It is in hundredth. A row with more than  $\text{iparm}[3]/100 \times \sqrt{n}$  nonzeros is treated as a dense node and is removed before ordering to save ordering time. Default is 1000 (10 actually).
- `iparm[4]`: metric for ordering method selection. Zero or a positive value means using estimated number of floating-point operations to select the best ordering method, while a negative value means using estimated number of factors. Default is zero (using floating-point operation number).
- `iparm[5]`: maximum supernode size. Each supernode can have at most `iparm[5]` rows. A supernode that has more than `iparm[5]` rows will be split into multiple smaller supernodes. Zero means that CKTSO determines the maximum supernode size according to the cache size. Negative one means no limitation for supernode size. Default is -1 (infinite).
- `iparm[6]`: minimum number of columns for supernode detection. A supernode must have at least `iparm[6]` columns. Default is 64.
- `iparm[7]`: whether to perform scaling. Scaling may improve the solution accuracy for some ill-conditioned matrices with the overhead of a small performance drop. Default is zero (scaling disabled).
- `iparm[8]`: whether the right-hand-side vector is highly sparse. Typically if the right-hand-side vector has less than 10% nonzeros, it can be regarded as "highly sparse". This parameter is used to control whether to use a faster substitution method. It is only effective in sequential column-mode solving. Default is zero (not highly sparse).

- `iparm[9]`: whether to control the number of threads for parallel factorization, refactorization, and solving. If enabled and the matrix is too small or too sparse, factorization and refactorization may use a single thread even if multiple threads have been created, and solving may use fewer threads than created. Default is 1 (thread number control enabled).
- `iparm[10]`: dynamic memory growth factor. It is in percentage. A larger value helps reduce reallocations at runtime but costs more memory usage. Default is 150 (1.5 actually).
- `iparm[11]`: initial number of rows for supernode creation. CKTSO allocates memory of `iparm[11]` rows when a new supernode is created. A larger value helps reduce reallocations at runtime but costs more memory usage. Default is 16.
- `iparm[12]`: static pivoting method. This parameter is effective only if `ax=NULL` when calling `CKTSO_Analyze` or `CKTSO_L_Analyze`. Zero means using the conventional matching-based static pivoting. A positive values means using a fill-in aware variant of the conventional method. A negative value means using a diagonal-first variant of the conventional method. Default is -1 (using the diagonal-first variant).
- `iparm[13]`: synchronization method for threads. Zero or a positive value means using blocked wait, while a negative value means using busy wait. The latter reduces threads' synchronization cost but consumes CPUs. Default is zero (using blocked wait).
- `iparm[14]`: timeout value for waiting for slave threads to exit, in milliseconds. When calling `CKTSO_DestroySolver` or `CKTSO_L_DestroySolver`, if one or more slave threads do not exit after `iparm[14]` milliseconds, the calling thread will continue. In this case, -13 (resource leak) will be returned. Default is -1 (infinite, waiting until all slave threads have exited).

**Please do not change other input parameters (after `iparm[14]`) which are not listed above.**

## 6.2 Output Parameters

- `oparm[0]`: time of matrix analysis, in microsecond (us), reported by `CKTSO_Analyze` and `CKTSO_L_Analyze`.
- `oparm[1]`: time of factorization or refactorization, in microsecond (us), reported by `CKTSO_Factorize`, `CKTSO_L_Factorize`, `CKTSO_Refactorize`, and `CKTSO_L_Refactorize`.
- `oparm[2]`: time of solving, in microsecond (us), reported by `CKTSO_Solve`, `CKTSO_L_Solve`, `CKTSO_SolveMV`, and `CKTSO_L_SolveMV`.
- `oparm[3]`: time of sorting factors, in microsecond (us), reported by

CKTSO\_SortFactors and CKTSO\_L\_SortFactors.

oparm[4]: number of off-diagonal pivots.

oparm[5]: number of nonzeros of L, including diagonal.

oparm[6]: number of nonzeros of U, excluding diagonal (U diagonal is one).  
           oparm[5] + oparm[6] is the number of the LU factors.

oparm[7]: number of supernodes.

oparm[8]: selected ordering method, from 1 to 8, corresponding to iparm[2].

oparm[9]: singular row index when -6 is returned.

oparm[10]: number of memory reallocations invoked, only reported in factorization.

oparm[11]: memory requirement in bytes when -4 is returned.

oparm[12]: current memory usage in bytes.

oparm[13]: maximum memory usage in bytes.

oparm[14]: number of rows completed with pivoting reduction, only reported after CKTSO\_Factorize or CKTSO\_L\_Factorize with fast factorization enabled. oparm[14]=n means that the entire matrix is completed with pivoting reduction, implying no change in the matrix structure.

oparm[15]: time of factorization-solve or refactorization-solve, in microsecond (us), reported by CKTSO\_FactorizeAndSolve, CKTSO\_L\_FactorizeAndSolve, CKTSO\_RefactorizeAndSolve, and CKTSO\_L\_RefactorizeAndSolve.

## 7. Using CKTSO Libraries

CKTSO is provided in the format of x64 dynamic-link libraries. Two individual libraries for 32-bit integers and 64-bit integers are provided. Link the correct library to the user's executable.

- On Linux, link CKTSO with "-L<library path> -lckts0" or "-L<library path> -lckts0\_1". Some additional libraries may also be needed. Specifically, add "-lpthread -lm -ldl" to the linking command if linking is not successful. On some Linux systems, "-lrt" is also required. Before running the executable, run "export LD\_LIBRARY\_PATH=<library path>" to set the library path if necessary.
- For Visual Studio on Windows, add "#pragma comment(lib, \"ckts0.lib\")" or "#pragma comment(lib, \"ckts0\_1.lib\")" to any place of the user's source code. cktso.lib or cktso\_1.lib is only required in linking. When running the executable, only cktso.dll or cktso\_1.dll is required. The \*.dll file should be put together with the executable or in the system directory (i.e., C:\Windows\ or C:\Windows\System32\).

## 8. Using CKTSO in Circuit Simulators

First, call `CKTSO_CreateSolver` to create a solver instance. After the matrix structure is constructed, call `CKTSO_Analyze`. Please note that `ax` can be `NULL` when calling `CKTSO_Analyze`. However, if `ax` is provided, the values specified to `CKTSO_Analyze` should have similar features of the matrix values during the iterations of the entire simulation process, or fill-ins may be dramatically increased, since `CKTSO_Analyze` determines a matrix ordering based on both the symbolic pattern and numerical values if `ax` is provided.

During the iterations of circuit simulation, call `CKTSO_Factorize` or `CKTSO_Refactorize` and `CKTSO_Solve` to solve a linear system in each iteration. It is crucial to correctly select `CKTSO_Factorize` and `CKTSO_Refactorize`. It is suggested that in DC simulation, `CKTSO_Factorize` should always be used. In transient simulation, the two functions can be selected based on the convergence situation. Intuitively, matrix values change more slowly when Newton-Raphson iterations are nearer convergence. Based on this observation, when Newton-Raphson iterations are converging, using `CKTSO_Refactorize` tends to be numerically stable. Conventional SPICE-style simulators use the following method to judge whether Newton-Raphson iterations are **converged**

$$|\mathbf{x}^{(k)} - \mathbf{x}^{(k-1)}| < \varepsilon_a + \varepsilon_r \times \min\{||\mathbf{x}^{(k)}||, ||\mathbf{x}^{(k-1)}||\}$$

where  $\varepsilon_a$  and  $\varepsilon_r$  are given absolute and relative thresholds, respectively. One can simply use the same method **with larger thresholds** to judge whether Newton-Raphson iterations are **converging**. If so, `CKTSO_Refactorize` can be used; otherwise `CKTSO_Factorize` should be used. The thresholds for judging whether Newton-Raphson iterations are converging may be empirically determined. Note that the first factorization must call `CKTSO_Factorize`.

`CKTSO_Factorize` has an argument to specify whether fast factorization based on pivoting reduction is enabled. Pivoting reduction may boost the performance and scalability of parallel factorization in most cases. However, in some extreme cases, it may degrade the overall circuit simulation performance. To avoid bad situations, fast factorization should not be always enabled. In a circuit simulation process, there are some independent simulation "*methods*". In DC simulation, direct Newton-Raphson iterations, GMIN stepping, source stepping, and pseudo-transient are popular DC simulation "*methods*". The entire transient simulation can be regarded as a single "*method*". The usage of fast factorization is recommended as follows. The first factorization of each "*method*" should disable fast factorization, while the other factorizations may enable fast factorization. The purpose of this strategy is to avoid inter-method influences. To be more conservative, for each different stepping value (e.g., each different GMIN value) in stepping-based methods (e.g., GMIN stepping), the first factorization may also disable fast factorization. In transient simulation, each time after the time node is rolled back due to divergence, the next

factorization should also disable fast factorization.

CKTSO\_Solve has an argument to specify whether a forced sequential solving will be used. It should also be carefully specified. Parallel solving needs a one-time scheduler initialization step which takes about 2-5X time of the sequential solving time. The scheduler initialization step needs to be called every time once the symbolic pattern of the LU factors changes. Thus, the following usage is recommended. In DC simulation, CKTSO\_Factorize and CKTSO\_Solve with forced sequential solving are recommended, as CKTSO\_Factorize changes the symbolic pattern of the LU factors frequently. In transient simulation, CKTSO\_Factorize and CKTSO\_Refactorize are selected based on the above-mentioned strategy, and whether to use forced sequential solving depends on the selection of CKTSO\_Factorize and CKTSO\_Refactorize. For the iterations which call CKTSO\_Factorize, solving should be forced sequential if  $oparm[14] < n$  (matrix structure changes), while for all other iterations, solving can be parallel. Please note that even if forced sequential solving is disabled, CKTSO may still use sequential solving, according to some internal strategies.