# User Guide for SuiteSparse:GraphBLAS

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VERSION 1.1.2, Dec 28, 2017

#### Abstract

SuiteSparse:GraphBLAS is a full implementation of the Graph-BLAS standard, which defines a set of sparse matrix operations on an extended algebra of semirings using an almost unlimited variety of operators and types. When applied to sparse adjacency matrices, these algebraic operations are equivalent to computations on graphs. GraphBLAS provides a powerful and expressive framework for creating graph algorithms based on the elegant mathematics of sparse matrix operations on a semiring.

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### 1 Introduction

The GraphBLAS standard defines sparse matrix and vector operations on an extended algebra of semirings. The operations are useful for creating a wide range of graph algorithms.

For example, consider the matrix-matrix multiplication,  $\mathbf{C} = \mathbf{AB}$ . Suppose  $\mathbf{A}$  and  $\mathbf{B}$  are sparse n-by-n Boolean adjacency matrices of two undirected graphs. If the matrix multiplication is redefined to use logical AND instead of scalar multiply, and if it uses the logical OR instead of add, then the matrix  $\mathbf{C}$  is the sparse Boolean adjacency matrix of a graph that has an edge (i,j) if node i in  $\mathbf{A}$  and node j in  $\mathbf{B}$  share any neighbor in common. The OR-AND pair forms an algebraic semiring, and many graph operations like this one can be succinctly represented by matrix operations with different semirings and different numerical types. GraphBLAS provides a wide range of built-in types and operators, and allows the user application to create new types and operators without needing to recompile the GraphBLAS library.

A full and precise definition of the GraphBLAS specification is provided in *The GraphBLAS C API Specification* by Aydın Buluç, Timothy Mattson, Scott McMillan, José Moreira, and Carl Yang [BMM+17], based on *GraphBLAS Mathematics* by Jeremy Kepner [Kep17]. The GraphBLAS C API Specification is available at http://graphblas.org. This version of SuiteSparse:GraphBLAS fully conforms to Version 1.1.0 of that specification. In this User Guide, aspects of the GraphBLAS specification that would be true for any GraphBLAS implementation are simply called "GraphBLAS." Details unique to this particular implementation are referred to as SuiteSparse:GraphBLAS.

SPEC: See the tag SPEC: for extensions to the spec. They are also placed in text boxes like this one. All functions and objects with a name of the form GxB\_\* and all macros with a name of the form GXB\_ are extensions to the spec. Functions and objects prefixed with GB\_ are internal to SuiteSparse:GraphBLAS and should not be referenced by user applications.

# 2 Basic Concepts

Since the *GraphBLAS C API Specification* provides a precise definition of GraphBLAS, not every detail of every function is provided here. For example, some error codes returned by GraphBLAS are self-explanatory, but since a specification must precisely define all possible error codes a function can return, these are listed in detail in the *GraphBLAS C API Specification*. However, including them here is not essential and the additional information on the page might detract from a clearer view of the essential features of the GraphBLAS functions.

This User Guide also assumes the reader is familiar with the MATLAB language, created by Cleve Moler. MATLAB supports only the conventional plus-times semiring on sparse double and complex matrices, but a MATLAB-like notation easily extends to the arbitrary semirings used in GraphBLAS. The matrix multiplication in the example in the Introduction can be written in MATLAB notation as C=A\*B, if the Boolean OR-AND semiring is understood. Relying on a MATLAB-like notation allows the description in this User Guide to be expressive, easy to understand, and terse at the same time. The GraphBLAS C API Specification also makes use of some MATLAB-like language, such as the colon notation.

MATLAB notation will always appear here in fixed-width font, such as C=A\*B(:,j). In standard mathematical notation it would be written as the matrix-vector multiplication  $\mathbf{C} = \mathbf{A}\mathbf{b}_j$  where  $\mathbf{b}_j$  is the jth column of the matrix  $\mathbf{B}$ . The GraphBLAS standard is a C API and SuiteSparse:GraphBLAS is written in C, and so a great deal of C syntax appears here as well, also in fixed-width font. This User Guide alternates between all three styles as needed.

# 2.1 Graphs and sparse matrices

Graphs can be huge, with many nodes and edges. A dense adjacency matrix for a graph of n nodes takes  $O(n^2)$  memory, which is impossible if n is, say, a million. Most graphs are sparse, however, with only O(n) edges, and graphs with millions of nodes and edges can easily be created by representing them as sparse matrices, where only explicit values need to be stored. In SuiteSparse:GraphBLAS, creating a million-by-million sparse matrix or a trillion-by-1 column vector can be done on quite easily on a commodity laptop, as long as it does not have too many explicit entries.

A sparse matrix data structure only stores a subset of the possible  $n^2$ entries, and it assumes the values of entries not stored have some implicit value. In conventional linear algebra, this implicit value is zero, but it differs with different semirings. Explicit values are called *entries* and they appear in the data structure. The pattern of a matrix defines where its explicit entries appear. It will be referenced in one of two equivalent ways. It can be viewed as a set of indices (i, j), where (i, j) is in the pattern of a matrix **A** if A(i,j) is an explicit value. It can also be viewed as a Boolean matrix S where S(i,j) is true if (i,j) is an explicit entry and false otherwise. In MATLAB notation, S=spones(A) or S=(A~=0), if the implicit value is zero. Later on in this User Guide, this pattern of A, however it is stored, is called A. pattern when used in MATLAB notation. The (i, j) pairs, and their values, can also be extracted from the matrix via the MATLAB expression [I,J,X] = find(A), where the kth tuple (I(k),J(k),X(k)) represents the explicit entry A(I(k), J(k)), with numerical value X(k) equal to  $a_{ij}$ , with row index i=I(k) and column index j=J(k).

The entries in the pattern of **A** can take on any value, including the implicit value, whatever it happens to be. This differs slightly from MAT-LAB, which always drops all explicit zeros from its sparse matrices. This is a minor difference but it cannot be done in GraphBLAS. For example, in the max-plus tropical algebra, the implicit value is negative infinity, and zero has a different meaning. Here, the MATLAB notation used will assume that no explicit entries are ever dropped because their explicit value happens to match the implicit value.

Graph Algorithms in the Language on Linear Algebra, Kepner and Gilbert, eds., provides a framework for understanding how graph algorithms can be expressed as matrix computations [KG11]. For additional background on sparse matrix algorithms, see also [Dav06] and [DRSL16].

# 2.2 Overview of GraphBLAS methods and operations

GraphBLAS provides a collection of *methods* to create, query, and free each of its nine different types of objects: sparse matrices, sparse vectors, types, operators (binary, unary and select), monoids, semirings, and a descriptor object used for parameter settings. Details are given in Section 4.

Once these objects are created they can be used in mathematical operations (not to be confused with the how the term operator is used in Graph-BLAS). A short summary of these operations and their nearest MATLAB analog is given in the table below.

operation	approximate MATLAB analog
matrix multiplication	C=A*B
element-wise operations	C=A+B and C=A.*B
reduction to a vector or scalar	s=sum(A)
apply unary operator	C=-A
transpose	C=A'
submatrix extraction	C=A(I,J)
submatrix assignment	C(I,J)=A

GraphBLAS can do far more than what MATLAB can do in these rough analogs, but the list provides a first step in describing what GraphBLAS can do. Details of each GraphBLAS operation are given in Section 5. With this brief overview, the full scope of GraphBLAS extensions of these operations can now be described.

GraphBLAS has 11 built-in scalar types: Boolean, single and double precision floating-point, and 8, 16, 32, and 64-bit signed and unsigned integers. In addition, user-defined scalar types can be created from nearly any C typedef, as long as the entire type fits in a fixed-size contiguous block of memory (of arbitrary size). All of these types can be used to create Graph-BLAS sparse matrices or vectors.

The scalar addition of conventional matrix multiplication is replaced with a monoid. A monoid is an associative and commutative binary operator z=f(x,y) where all three domains are the same (the types of x, y, and z), and where the operator has an identity value id such that f(x,id)=f(id,x)=x. Performing matrix multiplication with a semiring uses a monoid in place of the "add" operator, scalar addition being just one of many possible monoids. The identity value of addition is zero, since x+0=0+x=x. GraphBLAS includes eight built-in operators suitable for use as a monoid: min (with an identity value of positive infinity), max (whose identity is negative infinity), add (identity is zero) multiply (with an identity of one), and four logical operators: AND, OR, exclusive-OR, and Boolean equality. User-created monoids can be defined with any associative and commutative operator that has an identity value.

Finally, a semiring can use any built-in or user-defined binary operator z=f(x,y) as its "multiply" operator, as long as the type of its output, z matches the type of the semiring's monoid. The user application can create any semiring based on any types, monoids, and multiply operators, as long

these few rules are followed.

Just considering built-in types and operators, GraphBLAS can perform C=A\*B in 960 unique semirings. With typecasting, any of these 960 semirings can be applied to matrices C, A, and B of any of the 11 types, in any combination. This gives  $960 \times 11^3 = 1,277,760$  possible kinds of sparse matrix multiplication supported by GraphBLAS, and this is counting just built-in types and operators. By contrast, MATLAB provides just two semirings for its sparse matrix multiplication C=A\*B: plus-times-double and plus-times-complex, not counting the typecasting that MATLAB does when multiplying a real matrix times a complex matrix. All of the 1.3 million forms of matrix multiplication methods in SuiteSparse:GraphBLAS are typically just as fast as computing C=A\*B in MATLAB using its own native sparse matrix multiplication methods, and sometimes faster.

A monoid can also be used in a reduction operation, like s=sum(A) in MATLAB. MATLAB provides the plus, times, min, and max reductions of a real or complex sparse matrix as s=sum(A), s=prod(A), s=min(A), and s=max(A), respectively. In GraphBLAS, any monoid can be used (min, max, plus, times, AND, OR, exclusive-OR, equality, or any user-defined monoid, on any user-defined type).

Element-wise operations are also expanded from what can be done in MATLAB. Consider matrix addition, C=A+B in MATLAB. The pattern of the result is the set union of the pattern of A and B. In GraphBLAS, any binary operator can be used in this set-union "addition." The operator is applied to entries in the intersection. Entries in A but not B, or visa-versa, are copied directly into C, without any application of the binary operator. The accumulator operation for  $\mathbf{Z} = \mathbf{C} \odot \mathbf{T}$  described in Section 2.3 is one example of this set-union application of an arbitrary binary operator.

Consider element-wise multiplication, C=A.\*B in MATLAB. The operator (multiply in this case) is applied to entries in the set intersection, and the pattern of C just this set intersection. Entries in A but not B, or visa-versa, do not appear in C. In GraphBLAS, any binary operator can be used in this manner, not must scalar multiplication. The difference between element-wise "add" and "multiply" is not the operators, but whether or not the pattern of the result is the set union or the set intersection. In both cases, the operator is only applied to the set intersection.

Finally, GraphBLAS includes a *non-blocking* mode where operations can be left pending, and saved for later. This is very useful for submatrix assignment (C(I,J)=A where I and J are integer vectors), or or scalar assignment

(C(i,j)=x where i and j are scalar integers). Because of how MATLAB stores its matrices, adding and deleting individual entries is very costly. For example, this is very slow in MATLAB, taking  $O(nz^2)$  time:

```
A = sparse (m,n);  % an empty sparse matrix
for k = 1:nz
    compute a value x, row index i, and column index j
    A (i,j) = x;
end
```

The above code is very easy read and simple to write, but exceedingly slow. In MATLAB, the method below is preferred and is far faster, taking only O(nz) time. It can easily be a million times faster than the method above. Unfortunately the second method below is a little harder to read and a little less natural to write:

GraphBLAS can do both methods. SuiteSparse:GraphBLAS stores its matrices in the same way as MATLAB except that it allows for pending computations, and as a result it can do both methods above equally as fast as the MATLAB sparse function, allowing the user to write simpler code.

#### 2.3 The accumulator and the mask

Most GraphBLAS operations can be modified via transposing input matrices, using an accumulator operator, applying a mask or its complement, and by clear all entries the matrix C after using it in the accumulator operator but before the final results are written back into it. All of these steps are optional, and are controlled by a descriptor object that holds parameter settings (see Section 4.9) that control the following options:

• the input matrices A and/or B can be transposed first.

- an accumulator operator can be used, like the plus in the statement C=C+A\*B. The accumulator operator can be any binary operator, and an element-wise "add" (set union) is performed using the operator.
- an optional mask can be used to selectively write the results to the output. The mask is a sparse Boolean matrix Mask whose size is the same size as the result. If Mask(i,j) is true, then the corresponding entry in the output can be modified by the computation. If Mask(i,j) is false, then the corresponding in the output is protected and cannot be modified by the computation. The Mask matrix acts exactly like logical matrix indexing in MATLAB, with one minor difference: in GraphBLAS notation, the mask operation is  $C\langle M \rangle = Z$ , where the mask M appears only on the left-hand side. In MATLAB, it would appear on both sides as C(Mask)=Z(Mask). If no mask is provided, the Mask matrix is implicitly all true. This is indicated by passing the value  $GrB_NULL$  in place of the Mask argument in GraphBLAS operations.

This process can be described in mathematical notation as:

```
\mathbf{A} = \mathbf{A}', if requested via descriptor (first input option)
```

 $\mathbf{B} = \mathbf{B}'$ , if requested via descriptor (second input option)

 ${f T}$  is computed according to the specific operation

 $\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot \mathbf{T}$ , accumulating and writing the results back via the mask

The application of the mask and the accumulator operator is written as  $\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot\mathbf{T}$  where  $\mathbf{Z} = \mathbf{C}\odot\mathbf{T}$  denotes the application of the accumulator operator, and  $\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{Z}$  denotes the mask operator via the Boolean matrix  $\mathbf{M}$ . The Accumulator Phase,  $\mathbf{Z} = \mathbf{C}\odot\mathbf{T}$ , is performed as follows:

Accumulator Phase: compute  $Z = C \odot T$ :

$$\begin{aligned} \mathbf{Z} &= \mathbf{T} \\ \text{else} \\ \mathbf{Z} &= \mathbf{C} \odot \mathbf{T} \end{aligned}$$

The accumulator operator is  $\odot$  in GraphBLAS notation, or accum in the code. The pattern of  $\mathbb{C} \odot \mathbb{T}$  is the set union of the patterns of  $\mathbb{C}$  and  $\mathbb{T}$ , and the operator is applied only on the set intersection of  $\mathbb{C}$  and  $\mathbb{T}$ . Entries in neither the pattern of  $\mathbb{C}$  nor  $\mathbb{T}$  do not appear in the pattern of  $\mathbb{Z}$ . That is:

```
for all entries (i, j) in \mathbf{C} \cap \mathbf{T} (that is, entries in both \mathbf{C} and \mathbf{T})
z_{ij} = c_{ij} \odot t_{ij}
for all entries (i, j) in \mathbf{C} \setminus \mathbf{T} (that is, entries in \mathbf{C} but not \mathbf{T})
z_{ij} = c_{ij}
for all entries (i, j) in \mathbf{T} \setminus \mathbf{C} (that is, entries in \mathbf{T} but not \mathbf{C})
z_{ij} = t_{ij}
```

The Accumulator Phase is followed by the Mask/Replace Phase,  $\mathbf{C}\langle\mathbf{M}\rangle=\mathbf{Z}$  as controlled by the Grb\_REPLACE and Grb\_SCMP descriptor options:

```
\begin{split} \mathbf{Mask/Replace\ Phase:\ compute\ } \mathbf{C}\langle\mathbf{M}\rangle &= \mathbf{Z}; \\ & \text{if } (\mathtt{GrB\_REPLACE}) \text{ delete all entries in } \mathbf{C} \\ & \text{if } \mathtt{Mask\ is\ NULL} \\ & \text{if } (\mathtt{GrB\_SCMP}) \\ & \mathbf{C} \text{ is not modified} \\ & \text{else} \\ & \mathbf{C} &= \mathbf{Z} \\ & \text{else} \\ & \text{if } (\mathtt{GrB\_SCMP}) \\ & \mathbf{C}\langle\neg\mathbf{M}\rangle &= \mathbf{Z} \\ & \text{else} \\ & \mathbf{C}\langle\mathbf{M}\rangle &= \mathbf{Z} \end{split}
```

Both phases of the accum/mask process are illustrated in MATLAB notation in Figure 1. A GraphBLAS operation starts with its primary computation, producing a result T; for matrix multiply, T=A\*B, or if A is transposed first, T=A'\*B, for example. Applying the accumulator, mask (or its complement) to obtain the final result matrix C can be expressed in the MATLAB accum\_mask function shown in the figure. This function is an exact, fully functional, and nearly-complete description of the GraphBLAS accumulator/mask operation. The only aspects it does not consider are typecasting (see Section 2.4), and the value of the implicit identity (for those, see another version in the Test folder).

One aspect of GraphBLAS cannot be as easily expressed in a MATLAB sparse matrix: namely, what is the implicit value of entries not in the pattern? To accommodate this difference in the accum\_mask MATLAB function, each sparse matrix A is represented with its values A.matrix and its pattern, A.pattern. The latter could be expressed as the sparse matrix A.pattern=spones(A) or A.pattern=(A~=0) in MATLAB, if the implicit

```
function C = accum_mask (C, Mask, accum, T, C_replace, Mask_complement)
[m n] = size (C.matrix) ;
Z.matrix = zeros (m, n) ;
Z.pattern = false (m, n) ;
if (isempty (accum))
   Z = T;
               % no accum operator
  % Z = accum (C,T), like Z=C+T but with an binary operator, accum
  p = C.pattern & T.pattern; Z.matrix (p) = accum (C.matrix (p), T.matrix (p));
  p = C.pattern & ~T.pattern; Z.matrix (p) = C.matrix (p);
  p = ~C.pattern & T.pattern ; Z.matrix (p) = T.matrix (p) ;
  Z.pattern = C.pattern | T.pattern ;
end
% = 1000 apply the mask to the values and pattern
C.matrix = mask (C.matrix, Mask, Z.matrix, C_replace, Mask_complement);
C.pattern = mask (C.pattern, Mask, Z.pattern, C_replace, Mask_complement) ;
end
function C = mask (C, Mask, Z, C_replace, Mask_complement)
% replace C if requested
if (C_replace)
   C(:,:) = 0;
if (isempty (Mask))
                                % if empty, Mask is implicit ones(m,n)
  % implicitly, Mask = ones (size (C))
   if (~Mask_complement)
     C = Z;
                                % this is the default
   else
                                % Z need never have been computed
     C = C;
   end
else
   % apply the mask
   if (~Mask_complement)
     C (Mask) = Z (Mask);
   else
     C (^{\sim}Mask) = Z (^{\sim}Mask);
   end
end
end
```

Figure 1: Applying the mask and accumulator,  $\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot \mathbf{T}$ 

value is zero. With different semirings, entries not in the pattern can be 1, +Inf, -Inf, or whatever is the identity value of the monoid. As a result, Figure 1 performs its computations on two MATLAB matrices: the values in A.matrix and the pattern in the logical matrix A.pattern. Implicit values are untouched.

The final computation in Figure 1 with a complemented Mask is easily expressed in MATLAB as C(~Mask)=Z(~Mask) but this is costly if Mask is very sparse (the typical case). It can be computed much faster in MATLAB without complementing the sparse Mask via:

```
R = Z; R (Mask) = C (Mask); C = R;
```

A set of MATLAB functions that precisely compute the  $\mathbf{C}\langle\mathbf{M}\rangle=\mathbf{C}\odot\mathbf{T}$  operation according to the full GraphBLAS specification is provided in Suite-Sparse:GraphBLAS as  $\mathtt{GB\_spec\_accum.m}$ , which computes  $\mathbf{Z}=\mathbf{C}\odot\mathbf{T}$ , and  $\mathtt{GB\_spec\_mask.m}$ , which computes  $\mathbf{C}\langle\mathbf{M}\rangle=\mathbf{Z}$ . SuiteSparse:GraphBLAS includes a complete list of  $\mathtt{GB\_spec\_*}$  functions that illustrate every GraphBLAS operation; these are discussed in in Section 5.1.

The methods in Figure 1 rely heavily on MATLAB's logical matrix indexing. For those unfamiliar with logical indexing in MATLAB, here is short summary. Logical matrix indexing in MATLAB is written as A(Mask) where A is any matrix and Mask is a logical matrix the same size as A. The expression x=A(Mask) produces a column vector x consisting of the entries of A where Mask is true. On the left-hand side, logical submatrix assignment A(Mask)=x does the opposite, copying the components of the vector x into the places in A where Mask is true. For example, to negate all values greater than 10 using logical indexing in MATLAB:

```
>> A = magic (4)
A =
    16
                    3
                          13
     5
           11
                   10
                           8
            7
                    6
                          12
      4
           14
                   15
>> A (A>10) =
                  A (A>10)
A =
   -16
             2
                    3
                         -13
                   10
      5
          -11
                           8
     9
            7
                    6
                         -12
          -14
                 -15
                           1
```

In MATLAB, logical indexing with a sparse matrix A and sparse logical matrix Mask is very efficient since MATLAB supports sparse logical matrices. The Mask operator in GraphBLAS works identically as sparse logical indexing in MATLAB, and is equally as fast (or faster) in SuiteSparse:GraphBLAS.

### 2.4 Typecasting

If an operator z=f(x) or z=f(x,y) is used with inputs that do not match its inputs x or y, or if its result z does not match the type of the matrix it is being stored into, then the values are typecasted. Typecasting in Graph-BLAS extends beyond just operators. Almost all GraphBLAS methods and operations are able to typecast their results, as needed.

If one type can be typecasted into the other, they are said to be *compatible*. All built-in types are compatible with each other. GraphBLAS cannot typecast user-defined types thus any user-defined type is only compatible with itself. When GraphBLAS requires inputs of a specific type, or when one type cannot be typecast to another, the GraphBLAS function returns an error code, Grb\_DOMAIN\_MISMATCH (refer to Section 3.4 for a complete list of error codes). Typecasting can only be done between built-in types, and it follows the rules of the C language (not MATLAB) wherever the rules of C are well-defined. In particular, a large integer outside the range of a smaller one is wrapped, modulo style. This differs from MATLAB.

However, unlike MATLAB, the C language specification states that the results of typecasting a float or double to an integer type is not always defined. In SuiteSparse:GraphBLAS, whenever C leaves the result undefined the rules used in MATLAB are followed. In particular +Inf converts to the largest integer value, -Inf converts to the smallest (zero for unsigned integers), and NaN converts to zero. Other than these special cases, Suite-Sparse:GraphBLAS trusts the C compiler for the rest of its typecasting.

Typecasting to bool is fully defined in the C language specification, even for NaN. The result is false if the value compares equal to zero, and true otherwise. Thus NaN converts to true.

**SPEC:** the GraphBLAS API Specification states that typecasting follows the rules of C. Yet C leaves some typecasting undefined. Suite-Sparse:GraphBLAS provides a precise definition for all typecasting as an extension to the spec.

### 2.5 Notation and list of GraphBLAS operations

As a summary of what GraphBLAS can do, the following table lists all Graph-BLAS operations. Upper case letters denote a matrix, and lower case letters are vectors. Let  $\mathbf{AB}$  denote the multiplication of two matrices over a semiring. The semiring can use any binary operator as its "multiply" operator, and any commutative and associative monoid as its "add" operator.

a 5	1 1	C(X)
GrB_mxm	matrix-matrix multiply	$\mathbf{C}\langle\mathbf{M} angle=\mathbf{C}\odot\mathbf{AB}$
GrB_vxm	vector-matrix multiply	$\mathbf{w}'\langle\mathbf{m}' angle=\mathbf{w}'\odot\mathbf{u}'\mathbf{A}$
GrB_mxv	matrix-vector multiply	$\mathbf{w}\langle\mathbf{m} angle=\mathbf{w}\odot\mathbf{A}\mathbf{u}$
GrB_eWiseMult	element-wise,	$\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot (\mathbf{A} \otimes \mathbf{B})$
	set union	$\mathbf{w}\langle\mathbf{m}\rangle = \mathbf{w}\odot(\mathbf{u}\otimes\mathbf{v})$
GrB_eWiseAdd	element-wise,	$\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot (\mathbf{A} \oplus \mathbf{B})$
	set intersection	$\mathbf{w}\langle\mathbf{m}\rangle = \mathbf{w}\odot(\mathbf{u}\oplus\mathbf{v})$
GrB_extract	extract submatrix	$\mathbf{C}\langle\mathbf{M} angle=\mathbf{C}\odot\mathbf{A}(\mathbf{I},\mathbf{J})$
		$\mathbf{w}\langle\mathbf{m}\rangle=\mathbf{w}\odot\mathbf{u}(\mathbf{i})$
GxB_subassign	assign submatrix	$\mathbf{C}(\mathbf{I},\mathbf{J})\langle\mathbf{M} angle = \mathbf{C}(\mathbf{I},\mathbf{J})\odot\mathbf{A}$
	(with submask for $\mathbf{C}(\mathbf{I}, \mathbf{J})$ )	$\mathbf{w}(\mathbf{i})\langle\mathbf{m} angle = \mathbf{w}(\mathbf{i})\odot\mathbf{u}$
GrB_assign	assign submatrix	$\mathbf{C}\langle\mathbf{M} angle(\mathbf{I},\mathbf{J})=\mathbf{C}(\mathbf{I},\mathbf{J})\odot\mathbf{A}$
	(with mask for $\mathbf{C}$ )	$\mathbf{w}\langle\mathbf{m} angle(\mathbf{i})=\mathbf{w}(\mathbf{i})\odot\mathbf{u}$
GrB_apply	apply unary operator	$\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot f(\mathbf{A})$
		$\mathbf{w}\langle\mathbf{m} angle = \mathbf{w}\odot f(\mathbf{u})$
GxB_select	apply select operator	$\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot f(\mathbf{A}, \mathbf{k})$
		$\mathbf{w}\langle\mathbf{m}\rangle = \mathbf{w}\odot f(\mathbf{u},\mathbf{k})$
GrB_reduce	reduce to vector	$\mathbf{w}\langle\mathbf{m}\rangle = \mathbf{w}\odot[\oplus_{j}\mathbf{A}(:,j)]$
	reduce to scalar	$s = s \odot [\oplus_{ij} \mathbf{A}(I,J)]$
GrB_transpose	transpose	$\mathbf{C}\langle\mathbf{M} angle=\mathbf{C}\odot\mathbf{A}'$

Each operation takes an optional  $GrB_Descriptor$  argument that modifies the operation. The input matrices A and B can be optionally transposed, the mask M can be complemented, and C can be cleared of its entries after it is used in  $C = C \odot T$  but before the  $C(M) = C \odot T$  assignment. Vectors are never transposed via the descriptor.

Let  $\mathbf{A} \oplus \mathbf{B}$  denote the element-wise operator that produces a set union pattern (like A+B in MATLAB). Any binary operator can be used this way in GraphBLAS, not just plus. Let  $\mathbf{A} \otimes \mathbf{B}$  denote the element-wise operator that produces a set intersection pattern (like A.\*B in MATLAB); any binary operator can be used this way, not just times.

Reduction of a matrix **A** to a vector reduces the *i*th row of **A** to a scalar  $w_i$ . This is like w=sum(A') since by default, MATLAB reduces down the columns, not across the rows. Since the input matrix can be optionally transposed, selecting this option obtains the behavior of sum in MATLAB.

# 3 GraphBLAS Context and Sequence

A user application that directly relies on GraphBLAS must include the GraphBLAS.h header file:

```
#include "GraphBLAS.h"
```

The GraphBLAS.h file defines functions, types, and macros prefixed with GrB\_, GRB\_, GxB\_, and GXB\_ that may be used in user applications. The prefixes GrB\_ and GRB\_ denote items that appear in the official GraphBLAS API C Specification. The prefixes GxB\_ and GXB\_ refer to SuiteSparse-specific extensions to the GraphBLAS API. Both may be used in user applications but be aware that items with prefixes GxB\_ and GXB\_ will not appear in other implementations of the GraphBLAS standard.

There are a few functions and objects prefixed with GB\_ that also appear in GraphBLAS.h, but all names with this prefix are internal to Suite-Sparse:GraphBLAS and must not be referenced by user applications. They are not documented in this User Guide, and no guarantee at all is made about them in future versions of this package. They can change or even be removed without notice. In addition, no contents of any GraphBLAS object (A, say) should be dereferenced with A->whatever. This content is opaque to end user applications and can change without notice in future versions of this package. These names and content are technically visible to end-user applications, but this is only to enable the creation and use of polymorphic functions via the \_Generic keyword in ANSI C11.

**SPEC:** The following macros are extensions to the spec.

The GraphBLAS.h file includes all the definitions required to use Graph-BLAS, including the following macros that can assist a user application in compiling and using GraphBLAS.

There are two version numbers associated with SuiteSparse:GraphBLAS: the version of the GraphBLAS API Specification it conforms to, and the version of the implementation itself. These can be used in the following manner in a user application:

```
#if GXB >= GXB_VERSION (2,0,3)
... use features in GraphBLAS specification 2.0.3 ...
#else
... only use features in early specifications
```

#### #endif

```
#if GXB_IMPLEMENTATION > GXB_VERSION (1,4,0)
... use features from version 1.4.0 of a specific GraphBLAS implementation
#endif
```

SuiteSparse:GraphBLAS also defines the following strings with #define. Refer to the GraphBLAS.h file for details.

- GXB\_ABOUT: describes this particular implementation, copyright, and URL.
- GXB\_DATE: the date of this implementation.
- GXB\_SPEC: describes the GraphBLAS specification
- GXB\_SPEC\_DATE: the date of the GraphBLAS specification
- GXB\_LICENSE: the license for this particular implementation

Finally, SuiteSparse:GraphBLAS gives itself a unique name of the form GXB\_SUITESPARSE\_GRAPHBLAS that the user application can use in #ifdef tests. This is helpful in case a particular implementation provides non-standard features that extend the GraphBLAS specification, such as additional predefined built-in operators, or if a GraphBLAS implementation does not yet fully implement all of the GraphBLAS specification. The Suite-Sparse:GraphBLAS name is provided in its GraphBLAS.h file as:

#### #define GXB\_SUITESPARSE\_GRAPHBLAS

For example, SuiteSparse:GraphBLAS predefines additional built-in operators not in the specification. If the user application wishes to use these in any GraphBLAS implementation, an #ifdef can control when they are used. Refer to the examples in the GraphBLAS/Demo folder.

As another example, the GraphBLAS API Specification states that an implementation need not define the order in which <code>GrB\_Matrix\_build</code> assembles duplicate tuples in its <code>[I,J,X]</code> input arrays. As a result, no particular ordering should be relied upon in general. However, SuiteSparse:GraphBLAS does guarantee an ordering, and this guarantee will be kept in future versions of SuiteSparse:GraphBLAS as well. Since not all implementations will ensure a particular ordering, the following can be used to exploit the ordering returned by SuiteSparse:GraphBLAS.

```
#ifdef GXB_SUITESPARSE_GRAPHBLAS
// duplicates in I, J, X assembled in a specific order;
// results are well-defined even if op is not associative.
GrB_Matrix_build (C, I, J, X, nvals, op);
#else
// duplicates in I, J, X assembled in no particular order;
// results are undefined if op is not associative.
GrB_Matrix_build (C, I, J, X, nvals, op);
#endif
```

The remainder of this section describes GraphBLAS functions that create, modify, and destroy the GraphBLAS context, or provide utility methods for dealing with errors:

GraphBLAS function	purpose	Section
GrB_init	start up GraphBLAS	3.1
GrB_wait	force completion of pending operations	3.2
<pre>GrB_Info</pre>	status code returned by GraphBLAS functions	3.3
GrB_error	get more details on the last error	3.4
<pre>GrB_finalize</pre>	finish GraphBLAS	3.5

## 3.1 GrB\_init: initialize GraphBLAS

GrB\_init must be called before any other GraphBLAS operation. It defines the mode that GraphBLAS will use: blocking or non-blocking. With blocking mode, all operations finish before returning to the user application. With non-blocking mode, operations can be left pending, and are computed only when needed. Non-blocking mode can be much faster than blocking mode, by many orders of magnitude in extreme cases. Blocking mode should be used only when debugging a user application. The mode cannot be changed once it is set by GrB\_init.

GraphBLAS objects are opaque to the user application. This allows GraphBLAS to postpone operations and then do them later in a more efficient manner by rearranging them and grouping them together. In non-blocking mode, the computations required to construct an opaque GraphBLAS object might not be finished when the GraphBLAS method or operation returns to the user. However, user-provided arrays are not opaque, and GraphBLAS methods and operations that read them (such as GrB\_Matrix\_build) or write to them (such as GrB\_Matrix\_extractTuples) always finish reading them, or creating them, when the method or operation returns to the user application.

In addition, all methods and operations that extract values from a Graph-BLAS object and return them into non-opaque user arrays always ensure that the computations for that object are completed when the method returns, namely: GrB\_\*\_nvals, GrB\_\*\_extractElement, GrB\_\*\_extractTuples, and GrB\_\*\_reduce (to scalar). These methods only ensure that the computations for a single object are completed. Use GrB\_wait to ensure that all computations are completed (see Section 3.2).

### 3.2 GrB\_wait: wait for pending operations to finish

```
GrB_Info GrB_wait ( ) ;  // finish all pending computations
```

GrB\_wait forces all pending operations to complete. Blocking mode acts as if GrB\_wait is called whenever a GraphBLAS method or operation returns to the user application.

Unless specific rules are followed, non-blocking mode can be unpredictable if user-defined functions have side effects or if they rely on global variables not under the control of GraphBLAS. Suppose the user application creates a user-defined operator that accesses a global variable. That operator is then used in a GraphBLAS operation, which is left pending. If the user application then changes the global variable before pending operations complete, the pending operations will be eventually computed with this different value.

Worse yet, a user-defined operator might be freed before it is needed to finish a pending operation. This causes undefined behavior.

For best results with GraphBLAS, user-defined functions should not have side effects, nor should they access global variables outside the control of GraphBLAS. This allows the non-blocking mode to be used at its fullest level of performance. However, both of these features can safely be used in user-defined functions if the following specific rules are followed.

• User-defined functions may be called in any order when used in a GraphBLAS operation. This order may change in non-obvious ways, even in the same GraphBLAS operation. For example, SuiteSparse:-GraphBLAS relies on two different algorithms for computing the matrix multiplication C = A'B. If C is small, or if A is a vector, then it does not transpose A explicitly, but uses dot-products between column vectors of A and B instead. Otherwise, it transposes A and uses a sequence of sparse-matrix-times-sparse-vector operations. MATLAB uses the same algorithm in its built-in sparse matrix multiplication, C=A\*B, also written by this author. A user application in GraphBLAS has no control over the decision on which algorithm is used, and the heuristic used to select the algorithm may change in the future. Other GraphBLAS implementations may use entirely different algorithms. The GrB\_wait function has no effect on this order and the user application should not rely on any particular order used in a specific implementation of GraphBLAS.

- User-defined functions are permitted to access global variables. However, if they do so, the global variables they rely on should not be changed if any GraphBLAS methods or operations are still pending, assuming GraphBLAS is executing in non-blocking mode (see Section 3.1). To ensure this, the user application must call GrB\_wait before changing any global variables relied upon by user-defined functions. Alternatively, computations can be forced to complete on selected matrices and vectors via GrB\_\*\_nvals, GrB\_\*\_extractElement, GrB\_\*\_extractTuples, and GrB\_\*\_reduce (to scalar) applied to selected matrices and vectors. The GrB\_\*\_nvals function is particularly well-suited for this purpose since it is otherwise an extremely light-weight computation in SuiteSparse:GraphBLAS.
- If any GraphBLAS methods or operations are still pending, freeing user-defined types, operators, monoids, semirings, vectors, matrices, or descriptors leads to undefined behavior. A user application must call GrB\_wait before freeing any user-defined object, if a pending operation relies on it, or by selective completion via, say, GrB\_\*\_nvals. Alternatively, if the user application is about to terminate GraphBLAS (see GrB\_finalize below), then all GraphBLAS objects may be freed in any order, without calling GrB\_wait. Pending computations will simply be abandoned.

GrB\_wait ensures that all computations are completed for all objects. For specific objects, GrB\_\*\_nvals, GrB\_\*\_extractElement, GrB\_\*\_extractTuples, and GrB\_\*\_reduce (to scalar) ensure that the pending operations are completed just for the matrix or vector they operate on. No other GraphBLAS method or operation guarantees the completion of pending computations, even though they may happen to do so in any particular implementation. In the current version, SuiteSparse:GraphBLAS exploits the non-blocking mode in the GrB\_\*\_setElement methods and the GrB\_assign and GxB\_subassign operations. Future versions of SuiteSparse:GraphBLAS may extend this to other methods and operations. Refer to the example at the end of Section 2.2.

### 3.3 GrB\_Info: status code returned by GraphBLAS

Each GraphBLAS method and operation returns its status to the caller as its return value, an enumerated type (an enum) called GrB\_Info. The first two values in the following table denote a successful status, the rest are error codes.

GrB_SUCCESS	the method or operation was successful		
GrB_NO_VALUE	A(i,j) requested but not there. Its value is		
	implicit.		
GrB_UNINITIALIZED_OBJECT	object has not been initialized		
GrB_INVALID_OBJECT	object is corrupted		
GrB_NULL_POINTER	input pointer is NULL		
GrB_INVALID_VALUE	generic error code; some value is bad		
<pre>GrB_INVALID_INDEX</pre>	a row or column index is out of bounds; for		
	indices passed as scalars, not in a list.		
GrB_DOMAIN_MISMATCH	object domains are not compatible		
GrB_DIMENSION_MISMATCH	matrix dimensions do not match		
GrB_OUTPUT_NOT_EMPTY	output matrix already has values in it		
GrB_OUT_OF_MEMORY	out of memory		
<pre>GrB_INDEX_OUT_OF_BOUNDS</pre>	a row or column index is out of bounds; for		
	indices in a list of indices.		
GrB_PANIC	unrecoverable error. SuiteSparse:GraphBLAS		
	never panics, however.		

Not all GraphBLAS methods or operations can return all status codes. According to the GraphBLAS specification, any GraphBLAS method or operation can return an out-of-memory condition, GrB\_OUT\_OF\_MEMORY, or a panic, GrB\_PANIC. These two errors, and the GrB\_INDEX\_OUT\_OF\_BOUNDS error, are called *execution errors*. The other errors are called *API* errors. An API error is detecting immediately, regardless of the blocking mode. The detection of an execution error may be deferred until the pending operations complete.

In the discussions of each method and operation in this User Guide, most of the obvious error code returns are not discussed. For example, if a required input is a NULL pointer, then GrB\_NULL\_POINTER is returned. Only error codes specific to the method or that require elaboration are discussed here. For a full list of the status codes that each GraphBLAS function can return, refer to *The GraphBLAS C API Specification* [BMM<sup>+</sup>17].

### 3.4 GrB\_error: get more details on the last error

```
const char *GrB_error ( ) ;  // return a string describing the last error
```

Each GraphBLAS method and operation returns a GrB\_Info error code. The GrB\_error function returns additional information on the error in a thread-safe null-terminated string. The string returned by GrB\_error is statically allocated in thread local storage and must not be freed or modified. The simplest way to use it is just to print it out, such as:

```
info = GrB_some_method_here (...);
if (info != GrB_SUCCESS)
{
    printf ("%s\n", GrB_error ());
}
```

SuiteSparse:GraphBLAS reports many helpful details. For example, if a row or column index is out of bounds, the report will state what those bounds are. If a matrix dimension is incorrect, the mismatching dimensions will be provided. GrB\_BinaryOp\_new and GrB\_UnaryOp\_new record the name the function passed to them, and GrB\_Type\_new records the name of its type parameter, and these are printed if the user-defined types and operators are used incorrectly. Refer to the output of the example programs in the Demo folder, which intentionally generate errors to illustrate the use of GrB\_error.

# 3.5 GrB\_finalize: finish GraphBLAS

GrB\_finalize must be called as the last GraphBLAS operation, even after all calls to GrB\_free. All GraphBLAS objects created by the user application should be freed first, before calling GrB\_finalize since GrB\_finalize will not free those objects.

In non-blocking mode, GraphBLAS may leave some computations as pending. These computations can be safely abandoned if the user application frees all GraphBLAS objects it has created and then calls GrB\_finalize. There is no need to call GrB\_wait in this case.

# 4 GraphBLAS Objects and their Methods

GraphBLAS defines nine different objects to represent matrices and vectors, their scalar data type (or domain), binary and unary operators on scalar types, operators for selecting entries from a matrix or vector, monoids, semirings, and a *descriptor* object used to specify optional parameters that modify the behavior of a GraphBLAS operation.

The GraphBLAS API makes a distinction between *methods* and *operations*. A method is a function that works on a GraphBLAS object, creating it, destroying it, or querying its contents. An operation (not to be confused with an operator) acts on matrices and/or vectors in a semiring.

GrB_Type	a scalar data type
<pre>GrB_UnaryOp</pre>	a unary operator $z = f(x)$ , where z and x are scalars
<pre>GrB_BinaryOp</pre>	a binary operator $z = f(x, y)$ , where z, x, and y are scalars
<pre>GxB_SelectOp</pre>	a select operator
<pre>GrB_Monoid</pre>	an associative and commutative binary operator
	and its identity value
<pre>GrB_Semiring</pre>	a monoid that defines the "plus" and a binary operator
	that defines the "multiply" for an algebraic semiring
<pre>GrB_Matrix</pre>	a 2D sparse matrix of any type
<pre>GrB_Vector</pre>	a 1D sparse column vector of any type
<pre>GrB_Descriptor</pre>	a collection of parameters that modify an operation

Each of these objects is implemented in C as an opaque handle, which is a pointer to a data structure held by GraphBLAS. User applications may not examine the content of the object directly; instead, they can pass the handle back to GraphBLAS which will do the work. Assigning one handle to another is valid but it does not make a copy of the underlying object.

GraphBLAS provides 11 built-in types and 253 built-in operators. With these, 44 unique monoids and 960 unique semirings can be constructed.

**SPEC:** SuiteSparse:GraphBLAS predefines all unique monoids and semirings that can be constructed from built-in types and operators, as an extension to the spec. They appear in GraphBLAS.h. The GxB\_SelectOp object is an extension to GraphBLAS.

## 4.1 The GraphBLAS type: GrB\_Type

A GraphBLAS GrB\_Type defines the type of scalar values that a matrix or vector contains, and the type of scalar operands for a unary or binary operator. There are eleven built-in types, and a user application can define any types of its own as well. The built-in types correspond to built-in types in C (#include <stdbool.h> and #include <stdint.h>), and the classes in MATLAB, as listed in the following table.

GraphBLAS	C type	MATLAB	description	range
$_{ m type}$		class		
GrB_BOOL	bool	logical	Boolean	true $(1)$ , false $(0)$
GrB_INT8	int8_t	int8	8-bit signed integer	-128 to 127
GrB_UINT8	uint8_t	uint8	8-bit unsigned integer	0 to 255
GrB_INT16	int16_t	int16	16-bit integer	$-2^{15}$ to $2^{15}-1$
GrB_UINT16	$uint16_t$	uint16	16-bit unsigned integer	0 to $2^{16} - 1$
GrB_INT32	int32_t	int32	32-bit integer	$-2^{31}$ to $2^{31}-1$
GrB_UINT32	$uint32_t$	uint32	32-bit unsigned integer	0 to $2^{32} - 1$
GrB_INT64	int64_t	int64	64-bit integer	$-2^{63}$ to $2^{63}-1$
GrB_UINT64	$uint64_t$	uint64	64-bit unsigned integer	0 to $2^{64} - 1$
GrB_FP32	float	single	32-bit IEEE 754	-Inf to +Inf
GrB_FP64	double	double	64-bit IEEE $754$	-Inf to +Inf

The user application can also define new types based on any typedef in the C language whose values are held in a contiguous region of memory. For example, a user-defined GrB\_Type could be created to hold any C struct whose content is self-contained. A C struct containing pointers might be problematic because GraphBLAS would not know to dereference the pointers to traverse the entire "scalar" entry, but this can be done if the objects referenced by these pointers are not moved. A user-defined complex type with real and imaginary types can be defined, or even a "scalar" type containing a fixed-sized dense matrix (see Section 4.1.1). The possibilities are endless. GraphBLAS can create and operate on sparse matrices and vectors in any of these types, including any user-defined ones. For user-defined types, GraphBLAS simply moves the data around itself (via memcpy), and then passes the values back to user-defined functions when it needs to do any computations on the type. The next sections describe the methods for the GrB\_Type object:

GrB_Type_new	create a user-defined type
<pre>GxB_Type_size</pre>	return the size of a type
<pre>GrB_Type_free</pre>	free a user-defined type

#### 4.1.1 GrB\_Type\_new: create a user-defined type

GrB\_Type\_new creates a new user-defined type. The type is a handle, or a pointer to an opaque object. The handle itself must not be NULL on input, but the content of the handle can be undefined. On output, the handle contains a pointer to a newly created type. The ctype parameter is peculiar because GrB\_Type\_new is a C macro, not an actual function. The ctype parameter is not a variable, but an actual type in C, either built-in or defined by a typedef. The only requirement on ctype is that sizeof(ctype) is valid in C, and that the type reside in a contiguous block of memory so that it can be moved with memcpy. For example, to create a user-defined type called Complex for double-precision complex values using the ANSI C11 double complex type, the following can be used. A complete example can be found in the usercomplex.c and usercomplex.h files in the Demo folder.

```
#include <math.h>
#include <complex.h>

GrB_Type Complex ;
GrB_Type_new (&Complex, double complex) ;
```

To demonstrate the flexibility of the GrB\_Type, consider a "scalar" consisting of 4-by-4 floating-point matrix and a string. This type might be useful for the 4-by-4 translation/rotation/scaling matrices that arise in computer graphics, along with a string containing a description or even a regular expression that can be parsed and executed in a user-defined operator. All that is required is a fixed-size type, where sizeof(ctype) is a constant.

```
typedef struct
{
    float stuff [4][4];
    char whatstuff [64];
}
wildtype;

GrB_Type WildType;
GrB_Type_new (&WildType, wildtype);
```

With this type a sparse matrix can be created in which each entry consists of a 4-by-4 dense matrix stuff and a 64-character string whatstuff. GraphBLAS treats this 4-by-4 as a "scalar." Any GraphBLAS method or operation that simply moves data can be used with this type without any further information from the user application. For example, entries of this type can be assigned to and extracted from a matrix or vector, and matrices containing this type can be transposed. A working example (wildtype.c in the Demo folder) creates matrices and multiplies them with a user-defined semiring with this type.

Performing arithmetic on matrices and vectors with user-defined types requires operators to be defined. For example, the user application can define its own type for complex numbers, but then transposing the matrix with GraphBLAS will not compute the complex conjugate transpose. This corresponds to the array transpose in MATLAB (C=A.') instead of the complex conjugate transpose (C=A'). To compute the complex conjugate transpose, the application would need to create a user-defined unary operator to conjugate a user-defined complex scalar, and then apply it to the matrix before or after the transpose, via GrB\_apply. An extensive set of complex operators are provided in the usercomplex.c example in the Demo folder, along with an include file, usercomplex.h, that is suitable for inclusion in any user application. Thus, while GraphBLAS does not include any complex types or operators, SuiteSparse:GraphBLAS provides them in two simple "user" files in the Demo folder.

Refer to Section 6.7 for more details on these two example user-defined types.

#### 4.1.2 GxB\_Type\_size: return the size of a type

This function acts just like sizeof(type) in the C language. For example GxB\_Type\_size (&s, GrB\_INT32) sets s to 4, the same as sizeof(int32\_t).

**SPEC:** The GxB\_Type\_size function is an extension to the spec.

#### 4.1.3 GrB\_Type\_free: free a user-defined type

GrB\_Type\_free frees a user-defined type. Either usage:

```
GrB_Type_free (&type) ;
GrB_free (&type) ;
```

frees the user-defined type and sets type to NULL. It safely does nothing if passed a NULL handle, or if type == NULL on input.

It is safe to attempt to free a built-in type. SuiteSparse:GraphBLAS silently ignores the request and returns GrB\_SUCCESS. A user-defined type should not be freed until all operations using the type are completed. Suite-Sparse:GraphBLAS attempts to detect this condition but it must query a freed object in its attempt. This is hazardous and not recommended. Operations on such objects whose type has been freed leads to undefined behavior.

It is safe to first free a type, and then a matrix of that type, but after the type is freed the matrix can no longer be used. The only safe thing that can be done with such a matrix is to free it.

Note the function signature of GrB\_Type\_free, above. It is illustrated with the generic name, GrB\_free. Any of the nine GraphBLAS objects can be freed with the single function, GrB\_free. Refer to Section 4.10 for more details.

GraphBLAS includes many such generic functions. When describing a specific variation, a function is described with its specific name in this User Guide (such as GrB\_Type\_free). When discussing features applicable to all specific forms, the generic name is used instead (such as GrB\_free).

# **4.2** GraphBLAS unary operators: GrB\_UnaryOp, z = f(x)

A unary operator is a scalar function of the form z = f(x). The domain (type) of z and x need not be the same.

There are six kinds of built-in unary operators: one, identity, additive inverse, absolute value, multiplicative inverse, and logical negation. In the notation in the table below, T is any of the 11 built-in types and is a place-holder for BOOL, INT8, UINT8, ... FP32, or FP64. For example, GrB\_AINV\_INT32 is a unary operator that computes z=-x for two values x and z of type GrB\_INT32.

The logical negation operator  $\mathtt{GrB\_LNOT}$  only works on Boolean types. The  $\mathtt{GxB\_LNOT\_}T$  functions operate on inputs of type T, implicitly typecasting their input to Boolean and returning result of type T, with a value 1 for true and 0 for false. The operators  $\mathtt{GxB\_LNOT\_BOOL}$  and  $\mathtt{GrB\_LNOT}$  are identical. Considering all combinations, there are thus 67 built-in unary operators ((6 kinds of operators)  $\times$  (11 types), and  $\mathtt{GrB\_LNOT}$ ).

GraphBLAS name	types (domains)	expression	description
		z = f(x)	
$\mathtt{GxB\_ONE\_}T$	$T \to T$	z = 1	one
${\tt GrB\_IDENTITY\_}T$	$T \to T$	z = x	identity
${\tt GrB\_AINV\_}T$	$T \to T$	z = -x	additive inverse
${\tt GxB\_ABS\_}T$	$T \to T$	z =  x	absolute value
${\tt GrB\_MINV\_}T$	$T \to T$	z = 1/x	multiplicative inverse
${\tt GxB\_LNOT\_}T$	$T \to T$	$z = \neg(x \neq 0)$	logical negation
GrB_LNOT	$\texttt{bool} \to \texttt{bool}$	$z = \neg x$	logical negation

**SPEC:**  $GxB_ONE_T$ ,  $GxB_ABS_T$  and  $GxB_LNOT_T$  are extensions to the spec.

Integer division by zero normally terminates an application, but this is avoided in SuiteSparse:GraphBLAS. For details, see the binary  $\mathtt{GrB\_DIV\_}T$  operators.

**SPEC:** The definition of integer division by zero is an extension to the spec.

The next sections define the following methods for the GrB\_UnaryOp object:

```
GrB_UnaryOp_new create a user-defined unary operator GxB_UnaryOp_ztype return the type of the output z for z=f(x) GxB_UnaryOp_xtype return the type of the input x for z=f(x) free a user-defined unary operator
```

#### 4.2.1 GrB\_UnaryOp\_new: create a user-defined unary operator

GrB\_UnaryOp\_new creates a new unary operator. The new operator is returned in the unaryop handle, which must not be NULL on input. On output, its contents contains a pointer to the new unary operator.

The two types xtype and ztype are the GraphBLAS types of the input x and output z of the user-defined function z = f(x). These types may be built-in types or user-defined types, in any combination. The two types need not be the same, but they must be previously defined before passing them to GrB\_UnaryOp\_new.

The function argument to GrB\_UnaryOp\_new is a pointer to a user-defined function with the following signature:

```
void (*f) (void *z, const void *x);
```

When the function f is called, the arguments z and x are passed as (void \*) pointers, but they will be pointers to values of the correct type, defined by ztype and xtype, respectively, when the operator was created. The pointers will be unique. That is, the user function is never called with pointers that point to the same space.

#### **4.2.2** GxB\_UnaryOp\_ztype: return the type of z

GxB\_UnaryOp\_ztype returns the ztype of the unary operator, which is the type of z in the function z = f(x).

**SPEC:** The GxB\_UnaryOp\_ztype function is an extension to the spec.

#### **4.2.3** GxB\_UnaryOp\_xtype: return the type of x

GxB\_UnaryOp\_xtype returns the xtype of the unary operator, which is the type of x in the function z = f(x).

**SPEC:** The GxB\_UnaryOp\_xtype function is an extension to the spec.

#### 4.2.4 GrB\_UnaryOp\_free: free a user-defined unary operator

GrB\_UnaryOp\_free frees a user-defined unary operator. Either usage:

```
GrB_UnaryOp_free (&unaryop) ;
GrB_free (&unaryop) ;
```

frees the unaryop and sets unaryop to NULL. It safely does nothing if passed a NULL handle, or if unaryop == NULL on input. It does nothing at all if passed a built-in unary operator.

# **4.3** GraphBLAS binary operators: GrB\_BinaryOp, z = f(x,y)

A binary operator is a scalar function of the form z = f(x, y). The types of z, x, and y need not be the same.

SuiteSparse:GraphBLAS has 17 kinds of built-in binary operators of the form  $T \times T \to T$  that work on all 11 of the built-in types, T, for a total of 187 binary operators of this form. These are listed in the table below. For each of these operators, all domains (types) of the three operands are the same. The six comparison operators and three logical operators all return a result one for true and zero for false, in the same domain T as their inputs. These six comparison operators are useful as "multiply" operators for creating semirings with non-Boolean monoids.

GraphBLAS	types (domains)	expression	description
name		z = f(x, y)	
${\tt GrB\_FIRST\_}T$	$T \times T \to T$	z = x	first argument
${\tt GrB\_SECOND\_}T$	$T\times T\to T$	z = y	second argument
${\tt GrB\_MIN\_}T$	$T\times T\to T$	$z = \min(x, y)$	minimum
${\tt GrB\_MAX\_}T$	$T\times T\to T$	$z = \max(x, y)$	maximum
${\tt GrB\_PLUS\_}T$	$T\times T\to T$	z = x + y	addition
${\tt GrB\_MINUS\_}T$	$T\times T\to T$	z = x - y	subtraction
${\tt GrB\_TIMES\_}T$	$T\times T\to T$	z = xy	multiplication
${\tt GrB\_DIV\_}T$	$T\times T\to T$	z = x/y	division
$\texttt{GxB\_ISEQ\_}T$	$T \times T \to T$	z = (x == y)	equal
${\tt GxB\_ISNE\_}T$	$T\times T\to T$	$z = (x \neq y)$	not equal
${\tt GxB\_ISGT\_}T$	$T\times T\to T$	z = (x > y)	greater than
${\tt GxB\_ISLT\_}T$	$T\times T\to T$	z = (x < y)	less than
${\tt GxB\_ISGE\_}T$	$T\times T\to T$	$z = (x \ge y)$	greater than or equal
${\tt GxB\_ISLE\_}T$	$T\times T\to T$	$z = (x \le y)$	less than or equal
$\mathtt{GxB\_LOR\_}T$	$T \times T \to T$	$z = (x \neq 0) \lor (y \neq 0)$	logical OR
${\tt GxB\_LAND\_}T$	$T\times T\to T$	$z = (x \neq 0) \land (y \neq 0)$	logical AND
${\tt GxB\_LXOR\_}T$	$T\times T\to T$	$z = (x \neq 0) \lor (y \neq 0)$	logical XOR

**SPEC:** The  $\texttt{GxB\_IS*}\_T$  operators and the Boolean  $\texttt{GxB\_L*}\_T$  are extensions to the spec.

Another set of six kinds of built-in comparison operators have the form  $T \times T \to bool$ . They are defined for all eleven built-in types, for a total of 66 binary operators. Note that when T is bool, the six operators give the same

results as the six GxB\_IS\*\_BOOL operators in the table above. These six comparison operators are useful as "multiply" operators for creating semirings with Boolean monoids.

GraphBLAS	types (domains)	expression	description
name		z = f(x, y)	
${\tt GrB\_EQ\_}T$	$T \times T  o \mathtt{bool}$	z = (x == y)	equal
${\tt GrB\_NE\_}T$	$T\times T\to \texttt{bool}$	$z = (x \neq y)$	not equal
${\tt GrB\_GT\_}T$	$T\times T\to \texttt{bool}$	z = (x > y)	greater than
${\tt GrB\_LT\_}T$	$T\times T\to \texttt{bool}$	z = (x < y)	less than
${\tt GrB\_GE\_}T$	$T\times T\to \texttt{bool}$	$z = (x \ge y)$	greater than or equal
${\tt GrB\_LE\_}T$	$T\times T\to \texttt{bool}$	$z = (x \le y)$	less than or equal

Finally, GraphBLAS has three built-in binary operators that operate purely in the Boolean domain. These three are identical to the GxB\_L\*\_BOOL operators described above, just with a shorter name.

GraphBLAS	types (domains)	expression	description
name		z = f(x, y)	
GrB_LOR	$\texttt{bool} \times \texttt{bool} \to \texttt{bool}$	$z = x \vee y$	logical OR
GrB_LAND	$\texttt{bool} \times \texttt{bool} \to \texttt{bool}$	$z = x \wedge y$	logical AND
GrB_LXOR	$\mathtt{bool} \times \mathtt{bool} \to \mathtt{bool}$	$z = x \veebar y$	logical XOR

This gives a total of 256 built-in binary operators listed in the tables above: 187 of the form  $T \times T \to T$ , 66 of the form  $T \times T \to \mathsf{bool}$ , and three purely Boolean. There are 240 unique operators since 16 of the 26 Boolean operators are redundant.

There are two sets of built-in comparison operators in SuiteSparse:Graph-BLAS, but they are not redundant. They are identical except for the type (domain) of their output, z. The  $\mathtt{GrB\_EQ\_}T$  and related operators compare their inputs of type T and produce a Boolean result of true or false. The  $\mathtt{GxB\_ISEQ\_}T$  and related operators do the same comparison and produce a result with same type T as their input operands, returning one for true or zero for false. The  $\mathtt{IS*}$  comparison operators are useful when combining comparisons with other non-Boolean operators. For example, a  $\mathtt{PLUS-ISEQ}$  semiring counts how many terms of the comparison are true. With this semiring, matrix multiplication  $\mathbf{C} = \mathbf{AB}$  for two weighted undirected graphs  $\mathbf{A}$  and  $\mathbf{B}$  computes  $c_{ij}$  as the number of edges node i and j have in common that have identical edge weights. Since the output type of the "multiplier" operator in a semiring must match the type of its monoid, the Boolean  $\mathtt{EQ}$  cannot be

combined with a non-Boolean PLUS monoid to perform this operation.

Likewise, SuiteSparse:GraphBLAS has two sets of logical OR, AND, and XOR operators. Without the  $\_T$  suffix, the three operators  $\texttt{GrB\_LOR}$ ,  $\texttt{GrB\_LAND}$ , and  $\texttt{GrB\_LXOR}$  operate purely in the Boolean domain, where all input and output types are  $\texttt{GrB\_BOOL}$ . The second set  $(\texttt{GxB\_LOR\_}T \texttt{GxB\_LAND\_}T$  and  $\texttt{GxB\_LXOR\_}T$ ) provides Boolean operators to all 11 domains, implicitly typecasting their inputs from type T to Boolean and returning a value of type T that is 1 for true or zero for false. The set of  $\texttt{GxB\_L*\_}T$  operators are useful since they can be combined with non-Boolean monoids in a semiring.

**SPEC:** The definition of integer division by zero is an extension to the spec.

Floating-point operations follow the IEEE 754 standard. Thus, computing x/0 for a floating-point x results in +Inf if x is positive, -Inf if x is negative, and NaN if x is zero. The application is not terminated. However, integer division by zero normally terminates an application. Suite-Sparse:GraphBLAS avoids this by adopting the same rules as MATLAB, which are analogous to how the IEEE standard handles floating-point division by zero. For integers, when x is positive, x/0 is the largest positive integer, for negative x it is the minimum integer, and 0/0 results in zero. For example, for an integer x of type  $GrB_INT32$ , 1/0 is  $2^{31} - 1$  and (-1)/0 is  $-2^{31}$ . Refer to Section 4.1 for a list of integer ranges.

The next sections define the following methods for the <code>GrB\_BinaryOp</code> object:

GrB_BinaryOp_new	create a user-defined binary operator
<pre>GxB_BinaryOp_ztype</pre>	return the type of the output z for $z = f(x, y)$
<pre>GxB_BinaryOp_xtype</pre>	return the type of the input $x$ for $z = f(x, y)$
<pre>GxB_BinaryOp_ytype</pre>	return the type of the input y for $z = f(x, y)$
<pre>GrB_BinaryOp_free</pre>	free a user-defined binary operator

## 4.3.1 GrB\_BinaryOp\_new: create a user-defined binary operator

```
GrB_Info GrB_BinaryOp_new
(
    GrB_BinaryOp *binaryop,
    void *function,
    const GrB_Type ztype,
    const GrB_Type xtype,
    const GrB_Type ytype
);
// handle for the new binary operator
// pointer to the binary function
// type of output z
// type of input x
// type of input y

// type
```

GrB\_BinaryOp\_new creates a new binary operator. The new operator is returned in the binaryop handle, which must not be NULL on input. On output, its contents contains a pointer to the new binary operator.

The three types xtype, ytype, and ztype are the GraphBLAS types of the inputs x and y, and output z of the user-defined function z = f(x, y). These types may be built-in types or user-defined types, in any combination. The three types need not be the same, but they must be previously defined before passing them to  $GrB_BinaryOp_new$ .

The final argument to GrB\_BinaryOp\_new is a pointer to a user-defined function with the following signature:

```
void (*f) (void *z, const void *x, const void *y);
```

When the function f is called, the arguments z, x, and y are passed as (void \*) pointers, but they will be pointers to values of the correct type, defined by ztype, xtype, and ytype, respectively, when the operator was created. The pointers will be unique. That is, the user function is never called with pointers that point to the same space.

## **4.3.2** GxB\_BinaryOp\_ztype: return the type of z

 $GxB_BinaryOp_ztype$  returns the ztype of the binary operator, which is the type of z in the function z = f(x, y).

**SPEC:** The GxB\_BinaryOp\_ztype function is an extension to the spec.

## **4.3.3** GxB\_BinaryOp\_xtype: return the type of x

GxB\_BinaryOp\_xtype returns the xtype of the binary operator, which is the type of x in the function z = f(x, y).

**SPEC:** The GxB\_BinaryOp\_xtype function is an extension to the spec.

## **4.3.4** GxB\_BinaryOp\_ytype: return the type of y

GxB\_BinaryOp\_ytype returns the ytype of the binary operator, which is the type of y in the function z = f(x, y).

**SPEC:** The GxB\_BinaryOp\_ytype function is an extension to the spec.

## 4.3.5 GrB\_BinaryOp\_free: free a user-defined binary operator

GrB\_BinaryOp\_free frees a user-defined binary operator. Either usage:

```
GrB_BinaryOp_free (&op) ;
GrB_free (&op) ;
```

frees the op and sets op to NULL. It safely does nothing if passed a NULL handle, or if op == NULL on input. It does nothing at all if passed a built-in binary operator.

# 4.4 GraphBLAS select operators: GxB\_SelectOp

A select operator is a scalar function of the form  $z = f(i, j, m, n, a_{ij}, k)$  that is applied to the entries  $a_{ij}$  of an m-by-n matrix. The domain (type) of z is always boolean. The domain (type) of  $a_{ij}$  can be any built-in or user-defined type, or it can be  $GrB_NULL$  if the operator is type-generic.

The GxB\_SelectOp operator is used by GxB\_select (see Section 5.12) to select entries from a matrix. Each entry A(i,j) is evaluated with the operator, which returns true if the entry is to be kept in the output, or false if it is not to appear in the output. The signature of the select function f are as follows:

There are five built-in select operators listed in the table below. For the first four operators, **k** is a pointer to a single scalar of type <code>int64\_t</code>. Each operator can be used on any type, including user-defined types. User-defined select operators can also be created.

GraphBLAS name	MATLAB	description
	analog	
GxB_TRIL	C=tril(A,k)	true for $A(i,j)$ if $(j-i) \le k$
GxB_TRIU	<pre>C=triu(A,k)</pre>	true for $A(i,j)$ if $(j-i) >= k$
GxB_DIAG	<pre>C=diag(A,k)</pre>	true for $A(i,j)$ if $(j-i) == k$
GxB_OFFDIAG	C=A-diag(A,k)	true for $A(i,j)$ if $(j-i) != k$
GxB_NONZERO	C=A(A~=0)	true if A(i,j) is nonzero

**SPEC:** GxB\_SelectOp and all built-in functions in the table above are extensions to the spec.

The built-in  $GxB_NONZERO$  select operator is unique in that it is a function of the value of the entry  $a_{ij}$ , but it is still type-generic. It does this by simply returning false all bits in the value are zero, or true otherwise. This gives

the proper result for any built-in type, since integer and floating-point zeros are represented this way. For user-defined types, the function returns the same thing. This action is well-defined but its suitability for any particular user-defined type must be determined according to how the user application defines the type, and what a value with all bits zero means for this type. Whatever it means, if the bits of a value with a user-defined type are all zero, the function returns false, and if any bit is one, the <code>GxB\_NONZERO</code> function returns true.

The next sections define the following methods for the GxB\_SelectOp object:

```
GxB\_SelectOp\_new create a user-defined select operator GxB\_SelectOp\_xtype return the type of the input x free a user-defined select operator
```

## 4.4.1 GxB\_SelectOp\_new: create a user-defined select operator

GxB\_SelectOp\_new creates a new select operator. The new operator is returned in the selectop handle, which must not be NULL on input. On output, its contents contains a pointer to the new select operator.

The function argument to  $GxB\_SelectOp\_new$  is a pointer to a user-defined function with the signature described on the prior page. Given the properties of an entry  $a_{ij}$  in an m-by-n matrix, the function should return true if the entry should be kept in the output of  $GxB\_select$ , or false if it should not appear in the output.

The type xtype is the GraphBLAS type of the input x of the user-defined function z = f(i, j, m, n, x, k). The type may be built-in or user-defined, or it may even be  $\mathtt{GrB\_NULL}$ . If the xtype is  $\mathtt{GrB\_NULL}$ , then  $\mathtt{GxB\_select}$  does not pass the value of  $x = a_{ij}$  to the select function, but passes  $\mathtt{GrB\_NULL}$  for the input x to the user-defined select function.

## **4.4.2** GxB\_SelectOp\_xtype: return the type of x

GxB\_SelectOp\_xtype returns the xtype of the select operator, which is the type of x in the function z = f(i, j, m, n, x, k). If the select operator is type-generic, xtype is returned as GrB\_NULL.

## 4.4.3 GxB\_SelectOp\_free: free a user-defined select operator

GxB\_SelectOp\_free frees a user-defined select operator. Either usage:

```
GxB_SelectOp_free (&selectop) ;
GrB_free (&selectop) ;
```

frees the selectop and sets selectop to NULL. It safely does nothing if passed a NULL handle, or if selectop == NULL on input. It does nothing at all if passed a built-in select operator.

# 4.5 GraphBLAS monoids: GrB\_Monoid

A monoid is defined on a single domain (that is, a single type), T. It consists of an associative binary operator z = f(x, y) whose three operands x, y, and z are all in this same domain T (that is  $T \times T \to T$ ). The associative operator must also have an identity element, or "zero" in this domain, such that f(x,0) = f(0,x) = 0. Recall that an associative operator f(x,y) is one for which the condition f(a,f(b,c)) = f(f(a,b),c) always holds. That is, operator can be applied in any order and the results remain the same.

Four kinds of built-in operators (MIN, MAX, PLUS, TIMES) can be used to form monoids for each of the ten non-Boolean built-in types, and 12 can be used for Boolean monoids, all of which are listed in the table below. This is a total of 52 valid monoids that can be constructed from built-in types and operators, although 8 of the 12 Boolean monoids are redundant (the four remaining being OR, AND, XOR, and EQ). There are thus a total of 44 unique monoids that can be constructed using built-in binary operators. Since the built-in monoids are also commutative, all of them can be used to create a semiring. Recall that a commutative operator f(x,y) is one for which the condition f(a,b) = f(b,a) always holds. That is, the two operands can be swapped and the results remain the same. One of the components of a semiring is a commutative monoid.

GraphBLAS	types (domains)	expression	identity
name		z = f(x, y)	
${ t GrB\_MIN\_T}$	$T \times T \to T$	$z = \min(x, y)$	$+\infty$
${ t GrB\_MAX\_T}$	$T \times T \to T$	$z = \max(x, y)$	$-\infty$
${ t GrB\_PLUS\_T}$	$T \times T \to T$	z = x + y	0
${\tt GrB\_TIMES\_}T$	$T\times T\to T$	z = xy	1
GrB_LOR, GxB_LOR_BOOL	$\texttt{bool} \times \texttt{bool} \to \texttt{bool}$	$z = x \vee y$	false
GrB_LAND, GxB_LAND_BOOL	$\mathtt{bool} \times \mathtt{bool} \to \mathtt{bool}$	$z = x \wedge y$	true
<pre>GrB_LXOR, GxB_LXOR_BOOL</pre>	$\mathtt{bool} \times \mathtt{bool} \to \mathtt{bool}$	$z = x \veebar y$	false
GrB_EQ_BOOL, GxB_ISEQ_BOOL	$\texttt{bool} \times \texttt{bool} \to \texttt{bool}$	z = (x == y)	true

The next sections define the following methods for the GrB\_Monoid object:

GrB_Monoid_new	create a monoid
<pre>GxB_Monoid_operator</pre>	return the monoid operator
<pre>GxB_Monoid_identity</pre>	return the monoid identity value
GrB_Monoid_free	free a monoid

**SPEC:** The predefined monoids are an extension to the spec.

#### 4.5.1 GrB\_Monoid\_new: create a monoid

GrB\_Monoid\_new creates a monoid. The operator, op, must be an associative binary operator, either built-in or user-defined.

In the definition above, <type> is a type-generic place-holder. For built-in types, it is the C type corresponding to the built-in type (see Section 4.1), such as bool, int32\_t, float, or double. In this case, identity is a const scalar value of the particular type, not a pointer. For user-defined types, <type> is void \*, and thus identity is a not a scalar itself but a void \* pointer to a memory location containing the identity value of the user-defined operator, op.

## **4.5.2** GxB\_Monoid\_operator: return the monoid operator

GxB\_Monoid\_operator returns the binary operator of the monoid.

**SPEC:** The GxB\_Monoid\_operator function is an extension to the spec.

## **4.5.3** GxB\_Monoid\_identity: return the monoid identity

GxB\_Monoid\_identity returns the identity value of the monoid. The void \* pointer, identity, must be non-NULL and must point to a memory space of size at least equal to the size of the type of the monoid. The type size can be obtained via GxB\_Monoid\_operator to return the monoid additive operator, then GxB\_BinaryOp\_ztype to obtain the ztype, followed by GxB\_Type\_size to get its size.

**SPEC:** The GxB\_Monoid\_identity function is an extension to the spec.

#### 4.5.4 GrB\_Monoid\_free: free a monoid

GrB\_Monoid\_frees frees a monoid. Either usage:

```
GrB_Monoid_free (&monoid) ;
GrB_free (&monoid) ;
```

frees the monoid and sets monoid to NULL. It safely does nothing if passed a NULL handle, or if monoid == NULL on input. It does nothing at all if passed a built-in monoid.

# 4.6 GraphBLAS semirings: GrB\_Semiring

A semiring defines all the operators required to define the multiplication of two sparse matrices in GraphBLAS,  $\mathbf{C} = \mathbf{AB}$ . The "add" operator is a commutative and associative monoid, and the binary "multiply" operator defines a function z = fmult(x,y) where the type of z matches the exactly with the monoid type. SuiteSparse:GraphBLAS includes 960 predefined built-in semirings, which are all those that can be constructed from built-in types and operators. The next sections define the following methods for the  $\mathtt{GrB\_Semiring}$  object:

```
GrB_Semiring_new create a semiring
GxB_Semiring_add return the additive monoid of a semiring
GxB_Semiring_multipliy return the binary operator of a semiring
GrB_Semiring_free free a semiring
```

## 4.6.1 GrB\_Semiring\_new: create a semiring

GrB\_Semiring\_new creates a new semiring, with add being the additive monoid and multiply being the binary "multiply" operator. In addition to the standard error cases, the function returns GrB\_DOMAIN\_MISMATCH if the output (ztype) domain of multiply does not match the domain of the add monoid.

Using built-in types and operators, 960 unique semirings can be built. This count excludes redundant Boolean operators (for example GrB\_TIMES\_BOOL and GxB\_LAND\_BOOL are different operators but they are redundant since they always return the same result).

- 680 semirings with a multiplier  $T \times T \to T$  where T is non-Boolean, from the complete cross product of:
  - 4 add monoids (MIN, MAX, PLUS, TIMES)

- 17 multiply operators (FIRST, SECOND, MIN, MAX, PLUS, MINUS, TIMES, DIV, ISEQ, ISNE, ISGT, ISLT, ISGE, ISLE, LOR, LAND, LXOR)
- 10 non-Boolean types, T
- 240 semirings with a comparison operator  $T \times T \to \mathsf{bool}$ , where T is non-Boolean, from the complete cross product of:
  - 4 Boolean add monoids (LAND, LOR, LXOR, EQ)
  - 6 multiply operators (EQ, NE, GT, LT, GE, LE)
  - 10 non-Boolean types, T
- 40 semirings with purely Boolean types, bool × bool → bool, from the complete cross product of:
  - 4 Boolean add monoids (LAND, LOR, LXOR, EQ)
  - 10 multiply operators (FIRST, SECOND, LOR, LAND, LXOR, EQ, GT, LT, GE, LE)

**SPEC:** SuiteSparse:GraphBLAS pre-defines all 960 semirings that can be constructed from built-in types and operators, as an extension to the spec.

#### 4.6.2 GxB\_Semiring\_add: return the additive monoid of a semiring

GxB\_Semiring\_add returns the additive monoid of a semiring.

**SPEC:** The GxB\_Semiring\_add function is an extension to the spec.

## 4.6.3 GxB\_Semiring\_multiply: return multiply operator of a semiring

```
GrB_Info GxB_Semiring_multiply // return multiply operator of a semiring (

GrB_BinaryOp *multiply, // returns multiply operator of the semiring const GrB_Semiring semiring // semiring to query

);
```

GxB\_Semiring\_multiply returns the binary multiplicative operator of a semiring.

 $\mathbf{SPEC:}$  The <code>GxB\_Semiring\_multiply</code> function is an extension to the spec.

## 4.6.4 GrB\_Semiring\_free: free a semiring

```
GrB_Info GrB_free // free a user-created semiring (
GrB_Semiring *semiring // handle of semiring to free );
```

GrB\_Semiring\_free frees a semiring. Either usage:

```
GrB_Semiring_free (&semiring) ;
GrB_free (&semiring) ;
```

frees the semiring and sets semiring to NULL. It safely does nothing if passed a NULL handle, or if semiring == NULL on input. It does nothing at all if passed a built-in semiring.

# 4.7 GraphBLAS vectors: GrB\_Vector

Many of the methods for GraphBLAS vectors require a row index or a size. Many methods for matrices require both a row and column index, or a row and column dimension. These are all integers of a specific type, GrB\_Index, which is defined in GraphBLAS.h as

```
typedef uint64_t GrB_Index ;
```

Row and column indices of an nrows-by-ncols matrix range from zero to the nrows-1 for the rows, and zero to ncols-1 for the columns. Indices are zero-based, like C, and not one-based, like MATLAB. In Suite-Sparse:GraphBLAS, the largest size permitted for any integer of GrB\_Index is  $2^{60}$ . If compiled for use in MATLAB, this maximum size is reduced to match the MATLAB maximum size, which is  $2^{48} - 1$ .

This section describes a set of methods that create, modify, query, and destroy a GraphBLAS sparse vector, GrB\_Vector:

```
GrB_Vector_new
                               create a vector
GrB_Vector_dup
                               copy a vector
                               clear a vector of all entries
GrB_Vector_clear
GrB_Vector_size
                               return the size of a vector
GrB_Vector_nvals
                               return the number of entries in a vector
GxB_Vector_type
                               return the type of a vector
GrB_Vector_build
                               build a vector from a set of tuples
GrB_Vector_setElement
                               add a single entry to a vector
GrB_Vector_extractElement
                               get a single entry from a vector
GrB_Vector_extractTuples
                               get all entries from a vector
GrB_Vector_free
                               free a vector
```

#### 4.7.1 GrB\_Vector\_new: create a vector

```
GrB_Info GrB_Vector_new // create a new vector with no entries

(
GrB_Vector *v, // handle of vector to create
const GrB_Type type, // type of vector to create
const GrB_Index n // vector dimension is n-by-1
);
```

 $GrB_Vector_new$  creates a new n-by-1 sparse vector with no entries in it, of the given type. This is analogous to MATLAB statement v = sparse (n,1), except that GraphBLAS can create sparse vectors any type. The pattern of the new vector is empty.

## 4.7.2 GrB\_Vector\_dup: copy a vector

GrB\_Vector\_dup makes a deep copy of a sparse vector, like w=u in MAT-LAB. In GraphBLAS, it is possible, and valid, to write the following:

Then w and u can be used interchangeably. However, only a pointer reference is made, and modifying one of them modifies both, and freeing one of them leaves the other as a dangling handle that should not be used. If two different vectors are needed, then this should be used instead:

```
GrB_Vector u, w ;
GrB_Vector_new (&u, GrB_FP64, n) ;
GrB_Vector_dup (&w, u) ;  // like w = u, but making a deep copy
```

Then w and u are two different vectors that currently have the same set of values, but they do not depend on each other. Modifying one has no effect on the other.

#### 4.7.3 GrB\_Vector\_clear: clear a vector of all entries

GrB\_Vector\_clear clears all entries from a vector. All values v(i) are now equal to the implicit value, depending on what semiring ring is used to perform computations on the vector. The pattern of v is empty, just as if it were created fresh with GrB\_Vector\_new. Analogous with v (:) = 0 in MATLAB. The type and dimension of v do not change. In SuiteSparse:GraphBLAS, any pending updates to the vector are discarded.

#### 4.7.4 GrB\_Vector\_size: return the size of a vector

GrB\_Vector\_size returns the size of a vector (the number of rows). Analogous to n = length(v) or n = size(v,1) in MATLAB.

#### 4.7.5 GrB\_Vector\_nvals: return the number of entries in a vector

GrB\_Vector\_nvals returns the number of entries in a vector. Roughly analogous to nvals = nnz(v) in MATLAB, except that the implicit value in GraphBLAS need not be zero and nnz (short for "number of nonzeros") in MATLAB is better described as "number of explicit entries" in GraphBLAS.

Forced completion: All computations for the vector v are guaranteed to be finished when GrB\_Vector\_nvals method returns. That is, it acts like an object-specific GrB\_wait for just this particular vector v, which is a side-effect useful in its own right. For example, suppose the computations required for v rely upon a user-defined operator that accesses a user-controlled global variable outside the scope or control of GraphBLAS. If the user-application needs to modify or free the variable, GrB\_Vector\_nvals can be used to force all pending operations for this vector v to complete. The user application can then safely modify the global variable. A call to GrB\_Vector\_nvals(&nvals,v) only ensures that the computations require to compute v are finished; other pending computations for other objects may remain. To ensure that all pending computations are complete for all GraphBLAS objects, use GrB\_wait instead.

## 4.7.6 GxB\_Vector\_type: return the type of a vector

GxB\_Vector\_type returns the type of a vector. Analogous to type = class (v) in MATLAB.

**SPEC:** The GxB\_Vector\_type function is an extension to the spec.

## 4.7.7 GrB\_Vector\_build: build a vector from a set of tuples

GrB\_Vector\_build constructs a sparse vector w from a set of tuples, I and X, each of length nvals. The vector w must have already been initialized with GrB\_Vector\_new, and it must have no entries in it before calling GrB\_Vector\_build.

This function is just like GrB\_Matrix\_build (see Section 4.8.8), except that it builds a sparse vector instead of a sparse matrix. For a description of what GrB\_Vector\_build does, refer to GrB\_Matrix\_build. For a vector, the list of column indices J in GrB\_Matrix\_build is implicitly a vector of length nvals all equal to zero. Otherwise the methods are identical.

**SPEC:** As an extension to the spec, results are defined even if dup is non-associative.

## 4.7.8 GrB\_Vector\_setElement: add a single entry to a vector

GrB\_Vector\_setElement sets a single entry in a vector, w(i) = x. The operation is exactly like setting a single entry in an n-by-1 matrix, A(i,0) = x, where the column index for a vector is implicitly j=0. For further details of this function, see GrB\_Matrix\_setElement in Section 4.8.9.

## 4.7.9 GrB\_Vector\_extractElement: get a single entry from a vector

GrB\_Vector\_extractElement extracts a single entry from a vector,  $\mathbf{x} = \mathbf{v}(\mathbf{i})$ . The method is identical to extracting a single entry  $\mathbf{x} = \mathbf{A}(\mathbf{i}, \mathbf{0})$  from an n-by-1 matrix, so further details of this method are discussed in Section 4.8.10, which discusses GrB\_Matrix\_extractElement. In this case, the column index is implicitly  $\mathbf{j} = \mathbf{0}$ .

**Forced completion:** All computations for the vector **v** are guaranteed to be finished when the method returns.

## 4.7.10 GrB\_Vector\_extractTuples: get all entries from a vector

GrB\_Vector\_extractTuples extracts all tuples from a sparse vector, analogous to [I,~,X] = find(v) in MATLAB. This function is identical to its GrB\_Matrix\_extractTuples counterpart, except that the array of column indices J does not appear in this function. Refer to Section 4.8.11 where further details of this function are described.

**Forced completion:** All computations for the vector **v** are guaranteed to be finished when the method returns.

#### 4.7.11 GrB\_Vector\_free: free a vector

GrB\_Vector\_free frees a vector. Either usage:

```
GrB_Vector_free (&v) ;
GrB_free (&v) ;
```

frees the vector v and sets v to NULL. It safely does nothing if passed a NULL handle, or if v == NULL on input. In SuiteSparse:GraphBLAS, any pending updates to the vector are abandoned.

# 4.8 GraphBLAS matrices: GrB\_Matrix

This section describes a set of methods that create, modify, query, and destroy a GraphBLAS sparse matrix, GrB\_Matrix:

```
GrB_Matrix_new
                              create a matrix
GrB_Matrix_dup
                              copy a matrix
GrB_Matrix_clear
                              clear a matrix of all entries
                              return the number of rows of a matrix
GrB_Matrix_nrows
                              return the number of columns of a matrix
GrB_Matrix_ncols
                              return the number of entries in a matrix
GrB_Matrix_nvals
GxB_Matrix_type
                              return the type of a matrix
                              build a matrix from a set of tuples
GrB_Matrix_build
GrB_Matrix_setElement
                              add a single entry to a matrix
                              get a single entry from a matrix
GrB_Matrix_extractElement
                              get all entries from a matrix
GrB_Matrix_extractTuples
GrB_Matrix_free
                              free a matrix
```

#### 4.8.1 GrB\_Matrix\_new: create a matrix

```
GrB_Info GrB_Matrix_new // create a new matrix with no entries

(
GrB_Matrix *A, // handle of matrix to create
const GrB_Type type, // type of matrix to create
const GrB_Index nrows, // matrix dimension is nrows-by-ncols
const GrB_Index ncols
);
```

GrB\_Matrix\_new creates a new nrows-by-ncols sparse matrix with no entries in it, of the given type. This is analogous to the MATLAB statement A = sparse (nrows, ncols), except that GraphBLAS can create sparse matrices of any type.

## 4.8.2 GrB\_Matrix\_dup: copy a matrix

GrB\_Matrix\_dup makes a deep copy of a sparse matrix, like C=A in MAT-LAB. In GraphBLAS, it is possible, and valid, to write the following:

Then C and A can be used interchangeably. However, only a pointer reference is made, and modifying one of them modifies both, and freeing one of them leaves the other as a dangling handle that should not be used. If two different matrices are needed, then this should be used instead:

```
GrB_Matrix A, C ;
GrB_Matrix_new (&A, GrB_FP64, n) ;
GrB_Matrix_dup (&C, A) ;  // like C = A, but making a deep copy
```

Then C and A are two different matrices that currently have the same set of values, but they do not depend on each other. Modifying one has no effect on the other.

#### 4.8.3 GrB Matrix clear: clear a matrix of all entries

```
GrB_Info GrB_Matrix_clear // clear a matrix of all entries;
( // type and dimensions remain unchanged
GrB_Matrix A // matrix to clear
);
```

GrB\_Matrix\_clear clears all entries from a matrix. All values A(i,j) are now equal to the implicit value, depending on what semiring ring is used to perform computations on the matrix. The pattern of A is empty, just as if it were created fresh with GrB\_Matrix\_new. Analogous with A (:,:) = 0 in MATLAB. The type and dimensions of A do not change. In SuiteSparse:Graph-BLAS, any pending updates to the matrix are discarded.

#### 4.8.4 GrB\_Matrix\_nrows: return the number of rows of a matrix

GrB\_Matrix\_nrows returns the number of rows of a matrix (nrow=size(A,1) in MATLAB).

#### 4.8.5 GrB\_Matrix\_ncols: return the number of columns of a matrix

GrB\_Matrix\_ncols returns the number of columns of a matrix (ncols=size(A,2) in MATLAB).

#### 4.8.6 GrB Matrix nyals: return the number of entries in a matrix

```
GrB_Info GrB_Matrix_nvals // get the number of entries in a matrix (

GrB_Index *nvals, // matrix has nvals entries const GrB_Matrix A // matrix to query

);
```

GrB\_Matrix\_nvals returns the number of entries in a matrix, like nnz(A) in MATLAB.

Forced completion: All computations for the matrix A are guaranteed to be finished when the method returns. That is, it acts like an object-specific GrB\_wait for just this particular matrix A. Other pending computations for other objects may remain. To ensure that all pending computations are complete for all GraphBLAS objects, used GrB\_wait instead.

#### 4.8.7 GxB\_Matrix\_type: return the type of a matrix

GxB\_Matrix\_type returns the type of a matrix, like type=class(A) in MATLAB.

**SPEC:** The GxB\_Matrix\_type function is an extension to the spec.

## 4.8.8 GrB\_Matrix\_build: build a matrix from a set of tuples

GrB\_Matrix\_build constructs a sparse matrix C from a set of tuples, I, J, and X, each of length nvals. The matrix C must have already been initialized with GrB\_Matrix\_new, and it must have no entries in it before calling GrB\_Matrix\_build. Thus the dimensions and type of C are not changed by this function, but are inherited from the prior call to GrB\_Matrix\_new or GrB\_matrix\_dup.

An error is returned (GrB\_INDEX\_OUT\_OF\_BOUNDS) if any row index in I is greater than or equal to the number of rows of C, or if any column index in J is greater than or equal to the number of columns of C

Any duplicate entries with identical indices are assembled using the binary dup operator provided on input. All three types (x, y, z for z=dup(x,y)) must be identical. The types of dup, C and X must all be compatible. See Section 2.4 regarding typecasting and compatibility). The values in X are typecasted, if needed, into the type of dup. Duplicates are then assembled into a matrix T of the same type as dup, using  $T(i,j) = dup \ (T \ (i,j), X \ (k))$ . After T is constructed, it is typecasted into the result C. That is, typecasting does not occur at the same time as the assembly of duplicates.

**SPEC:** As an extension to the spec, results are defined even if dup is non-associative.

The GraphBLAS API Specification requires dup to be associative so that entries can be assembled in any order, and states that the result is undefined if dup is not associative. However, SuiteSparse:GraphBLAS guarantees a well-defined order of assembly. Entries in the tuples [I,J,X] are first sorted in increasing order of row and column index, with ties broken by the position of the tuple in the [I,J,X] list. If duplicates appear, they are assembled in the order they appear in the [I,J,X] input. That is, if the same

indices i and j appear in positions k1, k2, k3, and k4 in [I,J,X], where k1 < k2 < k3 < k4, then the following operations will occur in order:

```
T (i,j) = X (k1) ;

T (i,j) = dup (T (i,j), X (k2)) ;

T (i,j) = dup (T (i,j), X (k3)) ;

T (i,j) = dup (T (i,j), X (k4)) ;
```

This is a well-defined order but the user should not depend upon it when using other GraphBLAS implementations since the GraphBLAS API specification does not require this ordering.

However, SuiteSparse:GraphBLAS guarantees this ordering, and with this well-defined order, several operators become very useful. In particular, the SECOND operator results in the last tuple overwriting the earlier ones. The FIRST operator means the value of the first tuple is used and the others are discarded.

The acronym dup is used here for the name of binary function used for assembling duplicates, but this should not be confused with the \_dup suffix in the name of the function GrB\_Matrix\_dup. The latter function does not apply any operator at all, nor any typecasting, but simply makes a pure deep copy of a matrix.

The parameter X is a pointer to any C equivalent built-in type, or a void \* pointer. The GrB\_Matrix\_build function uses the \_Generic feature of ANSI C11 to detect the type of pointer passed as the parameter X. If X is a pointer to a built-in type, then the function can do the right typecasting. If X is a void \* pointer, then it can only assume X to be a pointer to a user-defined type that is the same user-defined type of C and dup. This function has no way of checking this condition that the void \* X pointer points to an array of the correct user-defined type, so behavior is undefined if the user breaks this condition.

The GrB\_Matrix\_build method is analogous to C = sparse (I,J,X) in MATLAB, with several important extensions that go beyond that which MATLAB can do. In particular, the MATLAB sparse function only provides one option for assembling duplicates (summation), and it can only build double, double complex, and logical sparse matrices.

## 4.8.9 GrB\_Matrix\_setElement: add a single entry to a matrix

 $GrB_Matrix_setElement$  sets a single entry in a matrix, C(i,j)=x. If the entry is already present in the pattern of C, it is overwritten with the new value. If the entry is not present, it is added to C. In either case, no entry is ever deleted by this function. Passing in a value of x=0 simply creates an explicit entry at position (i,j) whose value is zero, even if the implicit value is assumed to be zero.

An error is returned (Grb\_INVALID\_INDEX) if the row index i is greater than or equal to the number of rows of C, or if the column index j is greater than or equal to the number of columns of C. Note that this error code differs from the same kind of condition in Grb\_Matrix\_build, which returns Grb\_INDEX\_OUT\_OF\_BOUNDS. This is because Grb\_INVALID\_INDEX is an API error, and is caught immediately even in non-blocking mode, whereas Grb\_INDEX\_OUT\_OF\_BOUNDS is an execution error whose detection may wait until the computation completes sometime later.

The scalar **x** is typecasted into the type of **C**. Any value can be passed to this function and its type will be detected, via the \_Generic feature of ANSI C11. For a user-defined type, **x** is a **void** \* pointer that points to a memory space holding a single entry of this user-defined type. This user-defined type must exactly match the user-defined type of **C** since no typecasting is done between user-defined types.

Performance considerations: SuiteSparse:GraphBLAS exploits the non-blocking mode to greatly improve the performance of this method. Refer to the example shown in Section 2.2. If the entry exists in the pattern already, it is updated right away and the work is not left pending. Otherwise, it is placed in a list of pending updates, and the later on the updates are done all at once, using the same algorithm used for GrB\_Matrix\_build. In other words, setElement in SuiteSparse:GraphBLAS builds its own internal list of tuples [I,J,X], and then calls GrB\_Matrix\_build whenever the matrix is

needed in another computation, or whenever GrB\_wait is called.

As a result, if calls to setElement are mixed with calls to most other methods and operations (even extractElement) then the pending updates are assembled right away, which will be slow. Performance will be good if many setElement updates are left pending, and performance will be poor if the updates are assembled frequently.

A few methods and operations can be intermixed with setElement, in particular, some forms of the GrB\_assign and GxB\_subassign operations are compatible with the pending updates from setElement. Sections 5.10 gives more details on which GxB\_subassign and GrB\_assign operations can be interleaved with calls to setElement without forcing updates to be assembled. Other methods that do not access the existing entries may also be done without forcing the updates to be assembled, namely GrB\_Matrix\_clear (which erases all pending updates), GrB\_Matrix\_free, GrB\_Matrix\_ncols, GrB\_Matrix\_nrows, GxB\_Matrix\_type, and of course GrB\_Matrix\_setElement itself. All other methods and operations cause the updates to be assembled. Future versions of SuiteSparse:GraphBLAS may extend this list.

See Section 6.3 for an example of how to use GrB\_Matrix\_setElement.

## 4.8.10 GrB\_Matrix\_extractElement: get a single entry from a matrix

GrB\_Matrix\_extractElement extracts a single entry from a matrix x=A(i,j). An error is returned (GrB\_INVALID\_INDEX) if the row index i is greater

than or equal to the number of rows of C, or if column index j is greater than or equal to the number of columns of C.

If the entry is not present then GraphBLAS does not know its value, since its value depends on the implicit value, which is the identity value of the additive monoid of the semiring. It is not a characteristic of the matrix itself, but of the semiring it is used in. A matrix can be used in any compatible semiring, and even a mixture of semirings, so the implicit value can change as the semiring changes.

As a result, if the entry is present, x=A(i,j) is performed and the scalar x is returned with this value. The method returns  $GrB\_SUCCESS$ . If the entry is not present, x is not modified, and  $GrB\_NO\_VALUE$  is returned to the caller. What this means is up to the caller.

The function knows the type of the pointer x, so it can do typecasting as needed, from the type of A into the type of x. User-defined types cannot be typecasted, so if A has a user-defined type then x must be a void \* pointer that points to a memory space the same size as a single scalar of the type of A.

Forced completion: All computations for the matrix A are guaranteed to be finished when the method returns. In in particular, this method causes all pending updates from GrB\_setElement, GrB\_assign, or GxB\_subassign to be assembled, so its use can have performance implications. Calls to this function should not be arbitrarily intermixed with calls to these other two functions. Everything will work correctly and results will be predictable, it will just be slow.

## 4.8.11 GrB\_Matrix\_extractTuples:get all entries from a matrix

GrB\_Matrix\_extractTuples extracts all the entries from the matrix A, returning them as a list of tuples, analogous to [I,J,X]=find(A) in MAT-LAB. Entries in the tuples [I,J,X] are unique. No pair of row and column indices (i,j) appears more than once.

The GraphBLAS API specification states the tuples can be returned in any order. SuiteSparse:GraphBLAS chooses to always return them in sorted order, first by column index (all tuples in column 0 appear first, then column 1, and so on), and then within each column the tuples are sorted by row index. SuiteSparse:GraphBLAS guarantees this ordering but this should

not be expected of all implementations of GraphBLAS since their internal representation may differ from that used by SuiteSparse:GraphBLAS.

The number of tuples in the matrix A is given by GrB\_Matrix\_nvals(&anvals,A). If anvals is larger than the size of the arrays (nvals in the parameter list), an error GrB\_INSUFFICIENT\_SIZE is returned, and no tuples are extracted. If nvals is larger than anvals, then only the first anvals entries in the arrays I J, and X are modified, containing all the tuples of A, and the rest of I J, and X are left unchanged. On output, nvals contains the number of tuples extracted.

**Forced completion:** All computations for the matrix **A** are guaranteed to be finished when the method returns.

#### 4.8.12 GrB\_Matrix\_free: free a matrix

GrB\_Matrix\_free frees a matrix. Either usage:

```
GrB_Matrix_free (&A) ;
GrB_free (&A) ;
```

frees the matrix A and sets A to NULL. It safely does nothing if passed a NULL handle, or if A == NULL on input. In SuiteSparse:GraphBLAS, any pending updates to the matrix are abandoned.

# 4.9 GraphBLAS descriptors: GrB\_Descriptor

A GraphBLAS descriptor modifies the behavior of a GraphBLAS operation (not a operator). GraphBLAS operations are described in Section 5, and all of them have a final parameter of a descriptor. If the descriptor is NULL, defaults are used. No GraphBLAS method (Section 4) is modified by a descriptor, and neither are any unary or binary operators.

```
SPEC: The Gxb_DEFAULT option is an extension to the spec.
```

In the current GraphBLAS API Specification, there are four different components in a descriptor. The access to these parameters and their values is governed by two enum types, GrB\_Desc\_Field and GrB\_Desc\_Value:

The internal representation is opaque to the user, but in this User Guide the four descriptor fields of a descriptor desc are illustrated as an array of four items, as described in the list below. The underlying implementation need not be an array:

• desc [GrB\_0UTP] is a parameter that modifies the output of a Graph-BLAS operation. Currently, there are two possible settings. In the default case, the output is not cleared, and  $\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{Z} = \mathbf{C}\odot\mathbf{T}$  is computed as-is, where  $\mathbf{T}$  is the results of the particular GraphBLAS operation.

In the non-default case,  $\mathbf{Z} = \mathbf{C} \odot \mathbf{T}$  is first computed, using the results of  $\mathbf{T}$  and the accumulator  $\odot$ . After this is done, if the  $\mathtt{GrB\_OUTP}$  descriptor field is set to  $\mathtt{GrB\_REPLACE}$ , then the output is cleared of its entries. Next, the assignment  $\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{Z}$  is performed.

• desc [GrB\_MASK] is a parameter that modifies the Mask, even if the mask is not present.

If this parameter is set to its default value, and if the mask is not present (Mask==NULL) then implicitly Mask(i,j)=1 for all i and j. If the mask is present then Mask(i,j)=1 means that C(i,j) is to be modified by the  $C\langle M \rangle = Z$  update. Otherwise, if Mask(i,j)=0, then C(i,j) is not modified, even if Z(i,j) is an entry with a different value; that value is simply discarded.

If the desc [GrB\_MASK] parameter is set to GrB\_SCMP, then the use of the mask is complemented. In this case, if the mask is not present (Mask==NULL) then implicitly Mask(i,j)=0 for all i and j. This means that none of C is modified and the entire computation of Z might as well have been skipped. That is, a complemented empty mask means no modifications are made to the output object at all, except perhaps to clear it in accordance with the GrB\_OUTP descriptor. With a complemented mask, if the mask is present then Mask(i,j)=0 means that C(i,j) is to be modified by the  $C\langle M \rangle = Z$  update. Otherwise, if Mask(i,j)=1, then C(i,j) is not modified, even if Z(i,j) is an entry with a different value; that value is simply discarded.

Using a parameter to complement the Mask is very useful because constructing the actual complement of a very sparse mask is impossible since it has too many entries. If the number of places in C that should be modified is very small, then use a sparse mask without complementing it. If the number of places in C that should be protected from modification is very small, then use a sparse mask to indicate those places, and use a descriptor GrB\_MASK that complements the use of the mask.

• desc [GrB\_INP0] and desc [GrB\_INP1] modify the use of the first and second input matrices A and B of the GraphBLAS operation.

If the desc [GrB\_INPO] is set to GrB\_TRAN, then A is transposed before using it in the operation. Likewise, if desc [GrB\_INP1] is set to

GrB\_TRAN, then the second input, typically called B, is transposed.

Vectors are never transposed via the descriptor. If a method's first parameter is a matrix and the second a vector, then desc [GrB\_INP0] modifies the matrix parameter and desc [GrB\_INP1] is ignored. If a method's first parameter is a vector and the second a matrix, then desc [GrB\_INP1] modifies the matrix parameter and desc [GrB\_INP0] is ignored.

To clarify this in each function, the inputs are labeled as first input: and second input: in the function signatures.

## 4.9.1 GrB\_Descriptor\_new: create a new descriptor

```
GrB_Info GrB_Descriptor_new  // create a new descriptor
(
    GrB_Descriptor *descriptor // handle of descriptor to create
);
```

GrB\_Descriptor\_new creates a new descriptor, with all fields set to their defaults (output is not replaced, mask is not complemented, and neither input matrix is transposed).

#### 4.9.2 GrB\_Descriptor\_set: set a parameter in a descriptor

GrB\_Descriptor\_set sets a descriptor field (GrB\_OUTP, GrB\_MASK, GrB\_INPO, or GrB\_INP1) to a particular value (GxB\_DEFAULT, GrB\_SCMP, GrB\_TRAN, or GrB\_REPLACE). In the current specification, the following settings can be made:

Descriptor field	Default	Non-default
GrB_OUTP	GxB_DEFAULT: The output matrix is	GrB_REPLACE: After computing
	not cleared. The operation computes	$\mathbf{Z} = \mathbf{C} \odot \mathbf{T}$ , the output $\mathbf{C}$ is
	$ \mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot\mathbf{T}.$	cleared of all entries. Then
		$\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{Z}$ is performed.
GrB_MASK	GxB_DEFAULT: The Mask is not com-	GrB_SCMP: The Mask is comple-
	plemented. Mask(i,j)=1 means the	mented. Mask(i,j)=0 means the
	value $C_{ij}$ can be modified by the op-	value $C_{ij}$ can be modified by the
	eration, while Mask(i,j)=0 means	operation, while Mask(i,j)=1
	the value $C_{ij}$ shall not be modified	means the value $C_{ij}$ shall not be
	by the operation.	modified by the operation.
GrB_INPO	GxB_DEFAULT: The first input is not	GrB_TRAN: The first input is
	transposed prior to using it in the	transposed prior to using it in
	operation.	the operation. Only matrices are
		transposed, never vectors.
GrB_INP1	GxB_DEFAULT: The second input is	GrB_TRAN: The second input is
	not transposed prior to using it in	transposed prior to using it in
	the operation.	the operation. Only matrices are
		transposed, never vectors.

## 4.9.3 GxB\_Descriptor\_get: get a parameter from a descriptor

GxB\_Descriptor\_get returns the value of a single field in a descriptor.

SPEC: The GxB\_Descriptor\_get function is an extension to the spec.

## 4.9.4 GrB\_Descriptor\_free: free a descriptor

GrB\_Descriptor\_free frees a descriptor. Either usage:

```
GrB_Descriptor_free (&descriptor) ;
GrB_free (&descriptor) ;
```

frees the descriptor and sets descriptor to NULL. It safely does nothing if passed a NULL handle, or if descriptor == NULL on input.

There are currently no predefined descriptors, but if these are added in the future, this function will do nothing if passed a built-in descriptor.

# 4.10 GrB\_free: free any GraphBLAS object

Each of the nine GraphBLAS objects has GrB\_\*\_new and GrB\_\*\_free methods that are specific to each object. They can also be accessed by a generic function, GrB\_free, that works for all nine objects. If G is any of the nine GraphBLAS objects, the statement

```
GrB_free (&G) ;
```

frees the object and sets the variable G to NULL. It is safe to pass in a NULL handle, or to free an object twice:

However, the following sequence of operations is not safe. The first two are valid but the last statement will lead to undefined behavior.

Some objects are predefined, such as the built-in types. If a user application attempts to free a built-in object, SuiteSparse:GraphBLAS will safely do nothing. In all cases, the GrB\_free function in SuiteSparse:GraphBLAS always returns GrB\_SUCCESS.

# 5 GraphBLAS Operations

The next sections define each of the GraphBLAS operations, also listed in the table below.

GrB_mxm	matrix-matrix multiply	$\mathbf{C}\langle\mathbf{M} angle=\mathbf{C}\odot\mathbf{AB}$
<pre>GrB_vxm</pre>	vector-matrix multiply	$\mathbf{w}'\langle\mathbf{m}' angle=\mathbf{w}'\odot\mathbf{u}'\mathbf{A}$
<pre>GrB_mxv</pre>	matrix-vector multiply	$\mathbf{w}\langle\mathbf{m}\rangle=\mathbf{w}\odot\mathbf{A}\mathbf{u}$
GrB_eWiseMult	element-wise,	$\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot (\mathbf{A} \otimes \mathbf{B})$
	set union	$\mathbf{w}\langle\mathbf{m}\rangle=\mathbf{w}\odot(\mathbf{u}\otimes\mathbf{v})$
GrB_eWiseAdd	element-wise,	$\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot(\mathbf{A}\oplus\mathbf{B})$
	set intersection	$\mathbf{w}\langle\mathbf{m}\rangle=\mathbf{w}\odot(\mathbf{u}\oplus\mathbf{v})$
GrB_extract	extract submatrix	$\mathbf{C}\langle\mathbf{M} angle=\mathbf{C}\odot\mathbf{A}(\mathbf{I},\mathbf{J})$
		$\mathbf{w}\langle\mathbf{m} angle=\mathbf{w}\odot\mathbf{u}(\mathbf{i})$
GxB_subassign	assign submatrix	$\mathbf{C}(\mathbf{I},\mathbf{J})\langle\mathbf{M} angle=\mathbf{C}(\mathbf{I},\mathbf{J})\odot\mathbf{A}$
	(with submask for $C(I, J)$ )	$\mathbf{w}(\mathbf{i})\langle\mathbf{m} angle = \mathbf{w}(\mathbf{i})\odot\mathbf{u}$
GrB_assign	assign submatrix	$\mathbf{C}\langle\mathbf{M} angle(\mathbf{I},\mathbf{J})=\mathbf{C}(\mathbf{I},\mathbf{J})\odot\mathbf{A}$
	(with submask for <b>C</b> )	/ ··· · \ (•)
	(WIGH SUBINASK TOL C)	$\mathbf{w}\langle\mathbf{m} angle(\mathbf{i})=\mathbf{w}(\mathbf{i})\odot\mathbf{u}$
GrB_apply	apply unary operator	$\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot f(\mathbf{A})$
GrB_apply		
GrB_apply GxB_select		$\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot f(\mathbf{A})$
	apply unary operator	$\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot f(\mathbf{A})$ $\mathbf{w}\langle \mathbf{m} \rangle = \mathbf{w} \odot f(\mathbf{u})$
	apply unary operator	$\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot f(\mathbf{A})$ $\mathbf{w}\langle \mathbf{m} \rangle = \mathbf{w} \odot f(\mathbf{u})$ $\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot f(\mathbf{A}, \mathbf{k})$
GxB_select	apply unary operator apply select operator	$\begin{aligned} \mathbf{C}\langle\mathbf{M}\rangle &= \mathbf{C}\odot f(\mathbf{A}) \\ \mathbf{w}\langle\mathbf{m}\rangle &= \mathbf{w}\odot f(\mathbf{u}) \\ \mathbf{C}\langle\mathbf{M}\rangle &= \mathbf{C}\odot f(\mathbf{A},\mathbf{k}) \\ \mathbf{w}\langle\mathbf{m}\rangle &= \mathbf{w}\odot f(\mathbf{u},\mathbf{k}) \end{aligned}$
GxB_select	apply unary operator  apply select operator  reduce to vector	$\begin{aligned} \mathbf{C}\langle\mathbf{M}\rangle &= \mathbf{C}\odot f(\mathbf{A}) \\ \mathbf{w}\langle\mathbf{m}\rangle &= \mathbf{w}\odot f(\mathbf{u}) \\ \mathbf{C}\langle\mathbf{M}\rangle &= \mathbf{C}\odot f(\mathbf{A},\mathbf{k}) \\ \mathbf{w}\langle\mathbf{m}\rangle &= \mathbf{w}\odot f(\mathbf{u},\mathbf{k}) \\ \mathbf{w}\langle\mathbf{m}\rangle &= \mathbf{w}\odot [\oplus_{j}\mathbf{A}(:,j)] \end{aligned}$

# 5.1 The GraphBLAS specification in MATLAB

SuiteSparse:GraphBLAS includes a MATLAB implementation of nearly the entire GraphBLAS specification, including all built-in types and operators. The typecasting rules and integer operator rules from GraphBLAS are implemented in MATLAB via mexFunctions that call the GraphBLAS routines in C. All other functions are written purely in MATLAB M-files, and are given names of the form GB\_spec\_\*. All of these MATLAB interfaces and M-file functions they are provided in the software distribution of Suite-Sparse:GraphBLAS. The purpose of this is two-fold:

- Illustration and documentation: MATLAB is so expressive, and so beautiful to read and write, that the GB\_spec\_\* functions read almost like the exact specifications from the GraphBLAS C API Specification. Excerpts and condensed versions of these functions have already been used to this point in the User Guide, such as Figure 1, and the subsequent sections rely on them as well. This is why the discussion here is not just relegated to an Appendix on testing; the reader can benefit from studying the GB\_spec\_\* functions to understand what a Graph-BLAS operation is computing. For example, GrB\_mxm (Section 5.2) includes a condensed and simplified version of GB\_spec\_mxm.
- Testing: Testing the C interface to SuiteSparse:GraphBLAS is a significant challenge since it supports so many different kinds of operations on a vast range of semirings. It is difficult to tell from looking at the result from a C function in GraphBLAS if the result is correct. Thus, each function has been written twice: once in a highly-optimized function in C, and again in a simple and elegant MATLAB function. The latter is almost a direct translation of all the mathematics behind the GraphBLAS API Specification, so it is much easier to visually inspect the GB\_spec\_\* version in MATLAB to ensure the correct mathematics are being computed.

The following functions are included in the SuiteSparse:GraphBLAS software distribution. Each has a name of the form GB\_spec\_\*, and each of them is a "mimic" of a corresponding C function in GraphBLAS. Not all functions in the C API have a corresponding mimic; in particular, many of the vector functions can be computed directly with the corresponding matrix version in the MATLAB implementations. A list of these files is shown below:

MATLAB GB_spec function	corresponding GraphBLAS	Section
	function or method	
GB_spec_accum.m	$\mathbf{Z} = \mathbf{C} \odot \mathbf{T}$	2.3
GB_spec_mask.m	$\mathbf{C}\langle\mathbf{M} angle=\mathbf{Z}$	2.3
GB_spec_accum_mask.m	$\mathbf{C}\langle\mathbf{M} angle=\mathbf{C}\odot\mathbf{T}$	2.3
GB_spec_Vector_extractElement.m	<pre>GrB_Vector_extractElement</pre>	4.7.9
GB_spec_build.m	GrB_Matrix_build	4.8.8
<pre>GB_spec_Matrix_extractElement.m</pre>	<pre>GrB_Matrix_extractElement</pre>	4.8.10
<pre>GB_spec_extractTuples.m</pre>	${\tt GrB\_Matrix\_extractTuples}$	4.8.11
GB_spec_mxm.m	GrB_mxm	5.2
GB_spec_vxm.m	GrB_vxm	5.3
GB_spec_mxv.m	GrB_mxv	5.4
GB_spec_eWiseMult_Vector.m	GrB_eWiseMult_Vector	5.5
<pre>GB_spec_eWiseMult_Matrix.m</pre>	<pre>GrB_eWiseMult_Matrix</pre>	5.5
${\tt GB\_spec\_eWiseAdd\_Vector.m}$	<pre>GrB_eWiseAdd_Vector</pre>	5.6
<pre>GB_spec_eWiseAdd_Matrix.m</pre>	<pre>GrB_eWiseAdd_Matrix</pre>	5.6
GB_spec_Vector_extract.m	GrB_Vector_extract	5.7.1
<pre>GB_spec_Matrix_extract.m</pre>	<pre>GrB_Matrix_extract</pre>	5.7.2
GB_spec_Col_extract.m	<pre>GrB_Col_extract</pre>	5.7.3
GB_spec_subassign.m	GxB_subassign	5.8
GB_spec_assign.m	GrB_assign	5.9
GB_spec_apply.m	GrB_apply	5.11
GB_spec_select.m	GxB_select	5.12
GB_spec_reduce_to_vector.m	GrB_reduce (to vector)	5.13.1
GB_spec_reduce_to_scalar.m	GrB_reduce (to scalar)	5.13.3
GB_spec_transpose.m	GrB_transpose	5.14
-		

Additional files are included for creating test problems and providing inputs to the above files, or supporting functions:

MATLAB GB_spec function	purpose
GB_spec_compare.m	Compares output of C and MATLAB functions
GB_spec_random.m	Generates a random matrix
GB_spec_op.m	MATLAB mimic of built-in operators
GB_spec_operator.m	Like GrB_*Op_new
GB_spec_opsall.m	List operators, types, and semirings
GB_spec_semiring.m	Like GrB_Semiring_new
GB_spec_descriptor.m	mimics a GraphBLAS descriptor
GB_spec_identity.m	returns the identity of a monoid
<pre>GB_spec_matrix.m</pre>	conforms a MATLAB sparse matrix to GraphBLAS
GB_define.m	creates draft of GraphBLAS.h

An intensive test suite has been written that generates test graphs in MATLAB, then computes the result in both the C version of the Suite-Sparse:GraphBLAS and in the MATLAB GB\_spec\_\* functions. Each C function in GraphBLAS has a direct mexFunction interface that allow the test suite in MATLAB to call both functions.

This approach has its limitations:

- matrix classes: MATLAB only supports sparse double, sparse double complex, and sparse logical matrices. MATLAB can represent all eleven GraphBLAS types as dense matrices, so in all these specification M-files, the matrices are either in dense format in the corresponding MATLAB class, or they are held as sparse double or sparse logical, and the actual GraphBLAS type is held with it as a string member of a MATLAB struct. To ensure the correct typecasting is computed, most of the MATLAB scripts work on dense matrices, not sparse ones. As a result, the MATLAB GB\_spec\_\* function are not meant for production use, but just for testing and illustration.
- integer operations: MATLAB and GraphBLAS handle integer operations differently. In MATLAB, an integer result outside the range of the integer is set to maximum or minimum integer. For example, int8(127)+1 is 127. This is useful for many computations such as image processing, but GraphBLAS follows the C rules instead, where integer values wrap, modulo style. For example, in GraphBLAS and in C, incrementing (int8\_t) 127 by one results in -128. Of course, an alternative would be for a MATLAB interface to create its own integer operators, each of which would follow the MATLAB integer rules of arithmetic. However, this would obscure the purpose of these GB\_spec\_\* and GB\_mex\_\* test functions, which is to test the C API of GraphBLAS. When the GB\_spec\_\* functions need to perform integer computations and typecasting, they call GraphBLAS to do the work, instead doing the work in MATLAB. This ensures that the GB\_spec\_\* functions obtain the same results as their GraphBLAS counterparts.
- elegance: to simplify testing, each MATLAB mexFunction interface a GraphBLAS function is a direct translation of the C API. For example, GB\_mex\_mxm is a direct interface to the GraphBLAS GrB\_mxm, even down the order of parameters. This approach abandons some of the potential features of MATLAB for creating elegant M-file interfaces in a

highly usable form, such as the ability to provide fewer parameters when optional parameters are not in use. These mexFunctions, as written, are not meant to be usable in a user application. They are not highly documented. They are meant to be fast, and direct, to accomplish the goal of testing SuiteSparse:GraphBLAS in MATLAB and comparing their results with the corresponding GB\_spec\_\* function. They are not recommended for use in general applications in MATLAB.

• generality: the MATLAB mexFunction interface needs to test the C API directly, so it must access content of SuiteSparse:GraphBLAS objects that are normally opaque to an end user application. As a result, these mexFunctions do not serve as a general interface to any conforming GraphBLAS implementation, but only to SuiteSparse:GraphBLAS.

In the MATLAB mimic functions, GB\_spec\_\*, a GraphBLAS matrix A is represented as a MATLAB struct with the following components:

- A.matrix: the values of the matrix. If A.matrix is a sparse double matrix, it holds a typecasted copy of the values of a GraphBLAS matrix, unless the GraphBLAS matrix is also double (GrB\_FP64).
- A.pattern: a logical matrix holding the pattern; A.pattern(i,j)=true if (i,j) is in the pattern of A, and false otherwise.
- A.class: the MATLAB class of the matrix corresponding to one of the eleven built-in types. Normally this is simply class(A.matrix).
- A.values: most of the GraphBLAS test mexFunctions return their result as a MATLAB sparse matrix, in the double class. This works well for all types except for the 64-bit integer types, since a double has about 54 bits of mantissa which is less than the 64 bits available in a long integer. To ensure no bits are lots, these values are also returned as a vector. This enables GB\_spec\_compare to ensure the test results are identical down to the very last bit, and not just to within roundoff error. Nearly all tests, even in double precision, check for perfect equality, not just for results accurate to within round-off error.

# 5.2 GrB\_mxm: matrix-matrix multiply

```
GrB_Info GrB_mxm
                                     // C<Mask> = accum (C, A*B)
                                    // input/output matrix for results
    GrB_Matrix C,
   const GrB_Matrix Mask,
                                    // optional mask for C, unused if NULL
   const GrB_BinaryOp accum,
                                    // optional accum for Z=accum(C,T)
    const GrB_Semiring semiring,
                                    // defines '+' and '*' for A*B
    const GrB_Matrix A,
                                    // first input: matrix A
    const GrB_Matrix B,
                                    // second input: matrix B
    const GrB_Descriptor desc
                                    // descriptor for C, Mask, A, and B
);
```

GrB\_mxm multiplies two sparse matrices A and B using the semiring. The input matrices A and B may be transposed according to the descriptor, desc (which may be NULL) and then typecasted to match the multiply operator of the semiring. Next, T=A\*B is computed on the semiring, precisely defined in the GB\_spec\_mxm.m script. The actual algorithm exploits sparsity and does not take  $O(n^3)$  time, but what computes is the following:

```
[m s] = size (A.matrix) ;
[s n] = size (B.matrix);
T.matrix = zeros (m, n, multiply.ztype) ;
T.pattern = zeros (m, n, 'logical') ;
T.matrix (:,:) = identity ;
                                       % the identity of the semiring's monoid
T.class = multiply.ztype ;
                                       % the ztype of the semiring's multiply op
A = cast (A.matrix, multiply.xtype); % the xtype of the semiring's multiply op
B = cast (B.matrix, multiply.ytype);
                                       % the ytype of the semiring's multiply op
for j = 1:n
   for i = 1:m
       for k = 1:s
           % T (i,j) += A (i,k) * B (k,j), using the semiring
            if (A.pattern (i,k) && B.pattern (k,j))
               z = multiply (A (i,k), B (k,j));
               T.matrix (i,j) = add (T.matrix (i,j), z);
               T.pattern(i,j) = true;
            end
        end
    end
end
```

Finally, T is typecasted into the type of C, and the results are written back into C via the accum and Mask,  $\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot\mathbf{T}$ . The latter step is reflected in the MATLAB function  $\mathtt{GB\_spec\_accum\_mask.m}$ , discussed in Section 2.3.

# 5.3 GrB\_vxm: vector-matrix multiply

```
GrB_Info GrB_vxm
                                    // w'<Mask> = accum (w, u'*A)
    GrB_Vector w,
                                    // input/output vector for results
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w,t)
                                    // defines '+' and '*' for u'*A
    const GrB_Semiring semiring,
    const GrB_Vector u,
                                    // first input: vector u
    const GrB_Matrix A,
                                    // second input: matrix A
    const GrB_Descriptor desc
                                    // descriptor for w, mask, and A
);
```

GrB\_vxm multiplies a row vector u' times a matrix A. The matrix A may be first transposed according to desc (as the second input, GrB\_INP1); the column vector u is never transposed via the descriptor. The inputs u and A are typecasted to match the xtype and ytype inputs, respectively, of the multiply operator of the semiring. Next, an intermediate column vector t=A'\*u is computed on the semiring using the same method as GrB\_mxm. Finally, the column vector t is typecasted from the ztype of the multiply operator of the semiring into the type of w, and the results are written back into w using the optional accumulator accum and mask.

The last step is  $\mathbf{w}\langle \mathbf{m} \rangle = \mathbf{w} \odot \mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices.

Performance considerations: Because of the way SuiteSparse:Graph-BLAS stores its matrices and vectors, GrB\_vxm with its default descriptor can be slower than GrB\_mxv with its default descriptor, when the vector u is very sparse. If the user application needs to use GrB\_vxm repeatedly with very sparse vectors u, it can be faster to work on the transpose of A instead.

If the matrix is symmetric, then u'\*A is the same as A'\*u, except that in this case the operands to the semiring's multiplier operator are reversed. This has no effect if the multiplier operator is commutative, but an adjustment would need to be made if it were not (such as replacing FIRST with SECOND, and GE with LE, for example).

Using the non-default GrB\_TRAN descriptor for A makes the GrB\_vxm operation equivalent to GrB\_mxv with its default descriptor (with the operands reversed in the multiplier, as well). The reverse is true as well; GrB\_mxv with GrB\_TRAN is the same as GrB\_vxm with a default descriptor.

The breadth-first search in Section 6.1 uses GrB\_mxv instead of GrB\_vxm, since the graph is symmetric and the multiplier (AND) is commutative.

# 5.4 GrB\_mxv: matrix-vector multiply

```
GrB_Info GrB_mxv
                                    // w<Mask> = accum (w, A*u)
    GrB_Vector w,
                                    // input/output vector for results
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w,t)
                                    // defines '+' and '*' for A*B
    const GrB_Semiring semiring,
    const GrB_Matrix A,
                                    // first input: matrix A
    const GrB_Vector u,
                                    // second input: vector u
    const GrB_Descriptor desc
                                    // descriptor for w, mask, and A
);
```

GrB\_mxv multiplies a matrix A times a column vector u. The matrix A may be first transposed according to desc (as the first input); the column vector u is never transposed via the descriptor. The inputs A and u are typecasted to match the xtype and ytype inputs, respectively, of the multiply operator of the semiring. Next, an intermediate column vector t=A\*u is computed on the semiring using the same method as GrB\_mxm. Finally, the column vector t is typecasted from the ztype of the multiply operator of the semiring into the type of w, and the results are written back into w using the optional accumulator accum and mask.

The last step is  $\mathbf{w}\langle \mathbf{m} \rangle = \mathbf{w} \odot \mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices.

Performance considerations: Refer to the discussion of GrB\_vxm. In SuiteSparse:GraphBLAS, GrB\_mxv is very efficient when u is sparse or dense, when the default descriptor is used. When u is very sparse and the default descriptor is used, GrB\_mxv is also very efficient.

When u is very sparse and GrB\_INPO is set to its non-default GrB\_TRAN, then this method is not efficient. If an application needs to perform A'\*u repeatedly where u is very sparse, then it can be faster to explicitly transpose A first, and then to use this function with its default descriptor on the transpose. That is, compute C=A' via GrB\_transpose, and then use this method to repeatedly compute C\*u, without selecting GrB\_TRAN.

# 5.5 GrB\_eWiseMult: element-wise operations, set intersection

Element-wise "multiplication" is shorthand for applying a binary operator element-wise on two matrices or vectors A and B, for all entries that appear in the set intersection of the patterns of A and B. This is like A.\*B for two sparse matrices in MATLAB, except that in GraphBLAS any binary operator can be used, not just multiplication.

The pattern of the result of the element-wise "multiplication" is exactly this set intersection. Entries in A but not B, or visa versa, do not appear in the result.

Let  $\otimes$  denote the binary operator to be used. The computation  $\mathbf{T} = \mathbf{A} \otimes \mathbf{B}$  is given below. Entries not in the intersection of  $\mathbf{A}$  and  $\mathbf{B}$  do not appear in the pattern of  $\mathbf{T}$ . That is:

for all entries 
$$(i, j)$$
 in  $\mathbf{A} \cap \mathbf{B}$   
 $t_{ij} = a_{ij} \otimes b_{ij}$ 

Depending on what kind of operator is used and what the implicit value is assumed to be, this can give the Hadamard product. This is the case for A.\*B in MATLAB since the implicit value is zero. However, computing a Hadamard product is not necessarily the goal of the eWiseMult operation. It simply applies any binary operator, built-in or user-defined, to the set intersection of A and B, and discards any entry outside this intersection. Its usefulness in a user's application does not depend upon it computing a Hadamard product in all cases. The operator need not be associative, commutative, nor have any particular property except for type compatibility with A and B, and the output matrix C.

The generic name for this operation is GrB\_eWiseMult, which can be used for both matrices and vectors.

### 5.5.1 GrB\_eWiseMult\_Vector: element-wise vector multiply

```
GrB_Info GrB_eWiseMult
                                    // w<Mask> = accum (w, u.*v)
    GrB_Vector w,
                                    // input/output vector for results
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w,t)
    const <operator> multiply,
                                    // defines '.*' for t=u.*v
    const GrB_Vector u,
                                    // first input: vector u
    const GrB_Vector v,
                                    // second input: vector v
    const GrB_Descriptor desc
                                    // descriptor for w and mask
);
```

GrB\_eWiseMult\_Vector computes the element-wise "multiplication" of two vectors  $\mathbf{u}$  and  $\mathbf{v}$ , element-wise using any binary operator (not just times). The vectors are not transposed via the descriptor. The vectors  $\mathbf{u}$  and  $\mathbf{v}$  are first typecasted into the first and second inputs of the multiply operator. Next, a column vector  $\mathbf{t}$  is computed, denoted  $\mathbf{t} = \mathbf{u} \otimes \mathbf{v}$ . The pattern of  $\mathbf{t}$  is the set intersection of  $\mathbf{u}$  and  $\mathbf{v}$ . The result  $\mathbf{t}$  has the type of the output ztype of the multiply operator.

The operator is typically a GrB\_BinaryOp, but the method is type-generic for this parameter. If given a monoid (GrB\_Monoid), the additive operator of the monoid is used as the multiply binary operator. If given a semiring (GrB\_Semiring), the multiply operator of the semiring is used as the multiply binary operator.

The next and final step is  $\mathbf{w}\langle\mathbf{m}\rangle = \mathbf{w}\odot\mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices. Note for all GraphBLAS operations, including this one, the accumulator  $\mathbf{w}\odot\mathbf{t}$  is always applied in a set union manner, even though  $\mathbf{t} = \mathbf{u}\otimes\mathbf{v}$  for this operation is applied in a set intersection manner.

### 5.5.2 GrB\_eWiseMult\_Matrix: element-wise matrix multiply

```
GrB_Info GrB_eWiseMult
                                    // C<Mask> = accum (C, A.*B)
    GrB_Matrix C,
                                    // input/output matrix for results
    const GrB_Matrix Mask,
                                    // optional mask for C, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for Z=accum(C,T)
    const <operator> multiply,
                                    // defines '.*' for T=A.*B
    const GrB_Matrix A,
                                    // first input: matrix A
    const GrB_Matrix B,
                                    // second input: matrix B
    const GrB_Descriptor desc
                                    // descriptor for C, Mask, A, and B
);
```

GrB\_eWiseMult\_Matrix computes the element-wise "multiplication" of two matrices A and B, element-wise using any binary operator (not just times). The input matrices may be transposed first, according to the descriptor desc. They are then typecasted into the first and second inputs of the multiply operator. Next, a matrix T is computed, denoted  $\mathbf{T} = \mathbf{A} \otimes \mathbf{B}$ . The pattern of T is the set intersection of A and B. The result T has the type of the output ztype of the multiply operator.

The multiply operator is typically a GrB\_BinaryOp, but the method is type-generic for this parameter. If given a monoid (GrB\_Monoid), the additive operator of the monoid is used as the multiply binary operator. If given a semiring (GrB\_Semiring), the multiply operator of the semiring is used as the multiply binary operator.

The operation can be expressed in MATLAB notation as:

```
[nrows, ncols] = size (A.matrix);
T.matrix = zeros (nrows, ncols, multiply.ztype);
T.class = multiply.ztype;
p = A.pattern & B.pattern;
A = cast (A.matrix (p), multiply.xtype);
B = cast (B.matrix (p), multiply.ytype);
T.matrix (p) = multiply (A, B);
T.pattern = p;
```

The final step is  $\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{C} \odot \mathbf{T}$ , as described in Section 2.3. Note for all GraphBLAS operations, including this one, the accumulator  $\mathbf{C} \odot \mathbf{T}$  is always applied in a set union manner, even though  $\mathbf{T} = \mathbf{A} \otimes \mathbf{B}$  for this operation is applied in a set intersection manner.

## 5.6 GrB\_eWiseAdd: element-wise operations, set union

Element-wise "addition" is shorthand for applying a binary operator element-wise on two matrices or vectors A and B, for all entries that appear in the set intersection of the patterns of A and B. This is like A+B for two sparse matrices in MATLAB, except that in GraphBLAS any binary operator can be used, not just addition. The pattern of the result of the element-wise "addition" is the set union of the pattern of A and B. Entries in neither in A nor in B do not appear in the result.

Let  $\oplus$  denote the binary operator to be used. The computation  $\mathbf{T} = \mathbf{A} \oplus \mathbf{B}$  is exactly the same as the computation with accumulator operator as described in Section 2.3. It acts like a sparse matrix addition, except that any operator can be used. The pattern of  $\mathbf{A} \oplus \mathbf{B}$  is the set union of the patterns of  $\mathbf{A}$  and  $\mathbf{B}$ , and the operator is applied only on the set intersection of  $\mathbf{A}$  and  $\mathbf{B}$ . Entries not in either the pattern of  $\mathbf{A}$  or  $\mathbf{B}$  do not appear in the pattern of  $\mathbf{T}$ . That is:

```
for all entries (i, j) in \mathbf{A} \cap \mathbf{B}

t_{ij} = a_{ij} \oplus b_{ij}

for all entries (i, j) in \mathbf{A} \setminus \mathbf{B}

t_{ij} = a_{ij}

for all entries (i, j) in \mathbf{B} \setminus \mathbf{A}

t_{ij} = b_{ij}
```

The only difference between element-wise "multiplication" ( $\mathbf{T} = \mathbf{A} \otimes \mathbf{B}$ ) and "addition" ( $\mathbf{T} = \mathbf{A} \oplus \mathbf{B}$ ) is the pattern of the result, and what happens to entries outside the intersection. With  $\otimes$  the pattern of  $\mathbf{T}$  is the intersection; with  $\oplus$  it is the set union. Entries outside the set intersection are dropped for  $\otimes$ , and kept for  $\oplus$ ; in both cases the operator is only applied to those (and only those) entries in the intersection. Any binary operator can be used interchangeably for either operation.

Element-wise operations do not operate on the implicit values, even implicitly, since the operations make no assumption about the semiring. As a result, the results can be different from MATLAB, which can always assume the implicit value is zero. For example, C=A-B is the conventional matrix subtraction in MATLAB. Computing A-B in GraphBLAS with eWiseAdd will apply the MINUS operator to the intersection, entries in A but not B will be unchanged and appear in C, and entries in neither A nor B do not appear in C. For these cases, the results matches the MATLAB C=A-B. Entries in B but not A do appear in C but they are not negated; they cannot be subtracted

from an implicit value in A. This is by design. If conventional matrix subtraction of two sparse matrices is required, and the implicit value is known to be zero, use GrB\_apply to negate the values in B, and then use eWiseAdd with the PLUS operator, to compute A+(-B).

The generic name for this operation is GrB\_eWiseAdd, which can be used for both matrices and vectors.

There is another minor difference in two variants of the element-wise functions. If given a semiring, the eWiseAdd functions use the binary operator of the semiring's monoid, while the eWiseMult functions use the multiplicative operator of the semiring.

#### 5.6.1 GrB\_eWiseAdd\_Vector: element-wise vector addition

```
// w<Mask> = accum (w, u+v)
GrB_Info GrB_eWiseAdd
    GrB_Vector w,
                                    // input/output vector for results
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w,t)
    const <operator> add,
                                    // defines '+' for t=u+v
    const GrB_Vector u,
                                    // first input: vector u
    const GrB_Vector v,
                                    // second input: vector v
    const GrB_Descriptor desc
                                    // descriptor for w and mask
);
```

GrB\_eWiseAdd\_Vector computes the element-wise "addition" of two vectors u and v, element-wise using any binary operator (not just plus). The vectors are not transposed via the descriptor. Entries in the intersection of u and v are first typecasted into the first and second inputs of the add operator. Next, a column vector t is computed, denoted  $t = u \oplus v$ . The pattern of t is the set union of u and v. The result t has the type of the output ztype of the add operator.

The add operator is typically a GrB\_BinaryOp, but the method is type-generic for this parameter. If given a monoid (GrB\_Monoid), the additive operator of the monoid is used as the add binary operator. If given a semiring (GrB\_Semiring), the additive operator of the monoid of the semiring is used as the add binary operator.

The final step is  $\mathbf{w}\langle \mathbf{m} \rangle = \mathbf{w} \odot \mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices.

#### 5.6.2 GrB eWiseAdd Matrix: element-wise matrix addition

```
GrB_Info GrB_eWiseAdd
                                    // C<Mask> = accum (C, A+B)
    GrB_Matrix C,
                                    // input/output matrix for results
    const GrB_Matrix Mask,
                                    // optional mask for C, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for Z=accum(C,T)
                                    // defines '+' for T=A+B
    const <operator> add,
    const GrB_Matrix A,
                                    // first input: matrix A
    const GrB_Matrix B,
                                    // second input: matrix B
    const GrB_Descriptor desc
                                    // descriptor for C, Mask, A, and B
);
```

 $GrB\_eWiseAdd\_Matrix$  computes the element-wise "addition" of two matrices A and B, element-wise using any binary operator (not just plus). The input matrices may be transposed first, according to the descriptor desc. Entries in the intersection then typecasted into the first and second inputs of the add operator. Next, a matrix T is computed, denoted  $T = A \oplus B$ . The pattern of T is the set union of A and B. The result T has the type of the output ztype of the add operator.

The add operator is typically a GrB\_BinaryOp, but the method is type-generic for this parameter. If given a monoid (GrB\_Monoid), the additive operator of the monoid is used as the add binary operator. If given a semiring (GrB\_Semiring), the additive operator of the monoid of the semiring is used as the add binary operator.

The operation can be expressed in MATLAB notation as:

```
[nrows, ncols] = size (A.matrix);
T.matrix = zeros (nrows, ncols, add.ztype);
p = A.pattern & B.pattern;
A = GB_mex_cast (A.matrix (p), add.xtype);
B = GB_mex_cast (B.matrix (p), add.ytype);
T.matrix (p) = add (A, B);
p = A.pattern & ~B.pattern; T.matrix (p) = cast (A.matrix (p), add.ztype);
p = ~A.pattern & B.pattern; T.matrix (p) = cast (B.matrix (p), add.ztype);
T.pattern = A.pattern | B.pattern;
T.class = add.ztype;
```

Except for when typecasting is performed, this is identical to how the accum operator is applied in Figure 1.

The final step is  $C(M) = C \odot T$ , as described in Section 2.3.

### 5.7 GrB\_extract: submatrix extraction

The GrB\_extract function is a generic name for three specific functions: GrB\_Vector\_extract, GrB\_Col\_extract, and GrB\_Matrix\_extract. The generic name appears in the function signature, but the specific function name is used when describing what each variation does.

### 5.7.1 GrB\_Vector\_extract: extract subvector from vector

```
GrB_Info GrB_extract
                                    // w<mask> = accum (w, u(I))
                                    // input/output vector for results
   GrB_Vector w,
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
   const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w,t)
    const GrB_Vector u,
                                    // first input: vector u
                                    // row indices
    const GrB_Index *I,
    const GrB_Index ni,
                                    // number of row indices
    const GrB_Descriptor desc
                                    // descriptor for w and mask
) ;
```

GrB\_Vector\_extract extracts a subvector from another vector, identical to t = u (I) in MATLAB where I is an integer vector of row indices. Refer to GrB\_Matrix\_extract for further details; vector extraction is the same as matrix extraction with n-by-1 matrices. To extract all rows of a vector, as in t = u (:) in MATLAB, use I = GrB\_ALL. The final step is  $\mathbf{w}\langle\mathbf{m}\rangle = \mathbf{w}\odot\mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices.

#### 5.7.2 GrB Matrix extract: extract submatrix from matrix

```
GrB_Info GrB_extract
                                    // C<Mask> = accum (C, A(I,J))
    GrB_Matrix C,
                                    // input/output matrix for results
    const GrB_Matrix Mask,
                                    // optional mask for C, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for Z=accum(C,T)
    const GrB_Matrix A,
                                    // first input: matrix A
    const GrB_Index *I,
                                    // row indices
    const GrB_Index ni,
                                    // number of row indices
                                    // column indices
    const GrB_Index *J,
    const GrB_Index nj,
                                    // number of column indices
    const GrB_Descriptor desc
                                    // descriptor for C, Mask, and A
);
```

GrB\_Matrix\_extract extracts a submatrix from another matrix, identical to T = A(I,J) in MATLAB where I and J are integer vectors of row and column indices, respectively, except that indices are zero-based in Graph-BLAS and one-based in MATLAB. The input matrix A may be transposed first, via the descriptor. The type of T and A are the same.

Entries outside A(I,J) are not accessed and do not take part in the computation. More precisely, assuming the matrix A is not transposed, the matrix T is defined as follows:

If duplicate indices are present in I or J, the above method defines the result in T. Duplicates result in the same values of A being copied into different places in T.

To extract all rows of a matrix, as in T = A (:,J) in MATLAB, use  $I = GrB\_ALL$  as the input argument. For all columns of a matrix, use  $J = GrB\_ALL$ . The final step is  $\mathbf{C}(\mathbf{M}) = \mathbf{C} \odot \mathbf{T}$ , as described in Section 2.3.

#### 5.7.3 GrB\_Col\_extract: extract column vector from matrix

```
GrB_Info GrB_extract
                                    // w<mask> = accum (w, A(I,j))
    GrB_Vector w,
                                    // input/output matrix for results
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w,t)
    const GrB_Matrix A,
                                    // first input: matrix A
    const GrB_Index *I,
                                    // row indices
    const GrB_Index ni,
                                    // number of row indices
    const GrB_Index j,
                                    // column index
    const GrB_Descriptor desc
                                    // descriptor for w, mask, and A
);
```

GrB\_Col\_extract extracts a subvector from a matrix, identical to t = A (I,j) in MATLAB where I is an integer vector of row indices and where j is a single column index. The input matrix A may be transposed first, via the descriptor, which results in the extraction of a single row j from the matrix A, the result of which is a column vector w. The type of t and A are the same. To extract all rows of a matrix, as in t = A (:,j) in MATLAB, use  $I = GrB_ALL$  as the input argument. The final step is  $\mathbf{w}\langle \mathbf{m} \rangle = \mathbf{w} \odot \mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices.

Performance considerations: Because of the way SuiteSparse:Graph-BLAS stores its matrices, row extraction is more costly than column extraction. That is, using the GrB\_TRAN option for the A matrix is slower than the default, which is to extract a column. If this function is to be used many times on the same matrix A to extract rows with the GrB\_TRAN option enabled, it can be faster to explicitly transpose the matrix A once, and then to extract columns instead.

## 5.8 GxB\_subassign: submatrix assignment

The methods described in this section are all variations of the form C(I, J)=A, which modifies a submatrix of the matrix C. All methods can be used in their generic form with the single name GxB\_subassign. This is reflected in the prototypes. However, to avoid confusion between the different kinds of assignment, the name of the specific function is used when describing each variation. If the discussion applies to all variations, the simple name GxB\_subassign is used.

GxB\_subassign is very similar to GrB\_assign, described in Section 5.9. The two operations are compared and contrasted in Section 5.10.

SPEC: All variants of GxB\_subassign are extensions to the spec.

### 5.8.1 GxB\_Vector\_subassign: assign to a subvector

```
GrB_Info GxB_subassign
                                    // w(I) < mask > = accum (w(I),u)
    GrB_Vector w,
                                    // input/output matrix for results
    const GrB_Vector mask,
                                    // optional mask for w(I), unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w(I),t)
    const GrB_Vector u,
                                    // first input: vector u
                                    // row indices
    const GrB_Index *I,
    const GrB_Index ni,
                                    // number of row indices
    const GrB_Descriptor desc
                                    // descriptor for w(I) and mask
);
```

GxB\_Vector\_subassign operates on a subvector w(I) of w, modifying it with the vector u. The method is identical to GxB\_Matrix\_subassign described in Section 5.8.2, where all matrices have a single column each. The mask has the same size as w(I) and u. The only other difference is that the input u in this method is not transposed via the GrB\_INPO descriptor.

### 5.8.2 GxB\_Matrix\_subassign: assign to a submatrix

```
GrB_Info GxB_subassign
                                    // C(I,J) < Mask > = accum (C(I,J),A)
    GrB_Matrix C,
                                    // input/output matrix for results
    const GrB_Matrix Mask,
                                    // optional mask for C(I,J), unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for Z=accum(C(I,J),T)
                                    // first input: matrix A
    const GrB_Matrix A,
    const GrB_Index *I,
                                    // row indices
                                    // number of row indices
    const GrB_Index ni,
    const GrB_Index *J,
                                    // column indices
    const GrB_Index nj,
                                    // number of column indices
    const GrB_Descriptor desc
                                    // descriptor for C(I,J), Mask, and A
);
```

GxB\_Matrix\_subassign operates only on a submatrix S of C, modifying it with the matrix A. For this operation, the result is not the entire matrix C, but a submatrix S=C(I,J) of C. The steps taken are as follows, except that A may be optionally transposed via the GrB\_INPO descriptor option.

Step	GraphBLAS	description
	notation	
1	S = C(I, J)	extract the $C(I, J)$ submatrix
2	$\mathbf{S}\langle\mathbf{M} angle=\mathbf{S}\odot\mathbf{A}$	apply the accumulator/mask to the submatrix ${f S}$
3	$\mathbf{C}(\mathbf{I},\mathbf{J}) = \mathbf{S}$	put the submatrix $S$ back into $C(I, J)$

The accumulator/mask step in Step 2 is the same as for all other Graph-BLAS operations, described in Section 2.3, except that for  $GxB\_subassign$ , it is applied to just the submatrix S = C(I, J), and thus the Mask has the same size as A, S, and C(I, J).

The GxB\_subassign operation is the reverse of matrix extraction:

- For submatrix extraction, GrB\_Matrix\_extract, the submatrix A(I,J) appears on the right-hand side of the assignment, C=A(I,J), and entries outside of the submatrix are not accessed and do not take part in the computation.
- For submatrix assignment, GxB\_Matrix\_subassign, the submatrix C(I, J) appears on the left-hand-side of the assignment, C(I, J)=A, and entries outside of the submatrix are not accessed and do not take part in the computation.

In both methods, the accumulator and mask modify the submatrix of the assignment; they simply differ on which side of the assignment the submatrix resides on. In both cases, if the Mask matrix is present it is the same size as the submatrix:

- For submatrix extraction,  $\mathbf{C}\langle\mathbf{M}\rangle=\mathbf{C}\odot\mathbf{A}(\mathbf{I},\mathbf{J})$  is computed, where the submatrix is on the right. The mask  $\mathbf{M}$  has the same size as the submatrix  $\mathbf{A}(\mathbf{I},\mathbf{J})$ .
- For submatrix assignment,  $\mathbf{C}(\mathbf{I}, \mathbf{J})\langle \mathbf{M} \rangle = \mathbf{C}(\mathbf{I}, \mathbf{J}) \odot \mathbf{A}$  is computed, where the submatrix is on the left. The mask  $\mathbf{M}$  has the same size as the submatrix  $\mathbf{C}(\mathbf{I}, \mathbf{J})$ .

In Step 1, the submatrix S is first computed by the GrB\_Matrix\_extract operation, S=C(I,J).

Step 2 accumulates the results  $S\langle M \rangle = S \odot T$ , exactly as described in Section 2.3, but operating on the submatrix S, not C, using the optional Mask and accum operator. The matrix T is simply T = A, or T = A' if A is transposed via the desc descriptor,  $GrB_INPO$ . The  $GrB_REPLACE$  option in the descriptor clears S after computing C = T or  $C = C \odot T$ , not all of C since this operation can only modify the specified submatrix of C.

Finally, Step 3 writes the result (which is the modified submatrix S and not all of C) back into the C matrix that contains it, via the assignment C(I,J)=S, using the reverse operation from the method described for matrix extraction:

Results are not defined for any GxB\_subassign operation if duplicate indices appear in I or J.

### 5.8.3 GxB\_Col\_subassign: assign to a sub-column of a matrix

```
GrB_Info GxB_subassign
                                    // C(I,j) < mask > = accum (C(I,j),u)
    GrB_Matrix C,
                                    // input/output matrix for results
                                    // optional mask for C(I,j), unused if NULL
    const GrB_Vector mask,
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(C(I,j),t)
    const GrB_Vector u,
                                    // input vector
    const GrB_Index *I,
                                    // row indices
                                    // number of row indices
    const GrB_Index ni,
    const GrB_Index j,
                                    // column index
    const GrB_Descriptor desc
                                    // descriptor for C(I,j) and mask
);
```

GxB\_Col\_subassign modifies a single sub-column of a matrix C. It is the same as GxB\_Matrix\_subassign where the index vector J[0]=j is a single column index (and thus nj=1), and where all matrices in GxB\_Matrix\_subassign (except C) consist of a single column. The mask vector has the same size as u and the sub-column C(I,j). The input descriptor GrB\_INPO is ignored; the input vector u is not transposed. Refer to GxB\_Matrix\_subassign for further details.

#### 5.8.4 GxB\_Row\_subassign: assign to a sub-row of a matrix

```
// C(i,J) < mask' > = accum (C(i,J),u')
GrB_Info GxB_subassign
    GrB_Matrix C,
                                    // input/output matrix for results
   const GrB_Vector mask,
                                    // optional mask for C(i,J), unused if NULL
   const GrB_BinaryOp accum,
                                    // optional accum for z=accum(C(i,J),t)
    const GrB_Vector u,
                                    // input vector
    const GrB_Index i,
                                    // row index
                                    // column indices
    const GrB_Index *J,
    const GrB_Index nj,
                                    // number of column indices
    const GrB_Descriptor desc
                                    // descriptor for C(i,J) and mask
);
```

GxB\_Row\_subassign modifies a single sub-row of a matrix C. It is the same as GxB\_Matrix\_subassign where the index vector I[0]=i is a single row index (and thus ni=1), and where all matrices in GxB\_Matrix\_subassign (except C) consist of a single row. The mask vector has the same size as u and the sub-column C(I,j). The input descriptor GrB\_INPO is ignored; the input vector u is not transposed. Refer to GxB\_Matrix\_subassign for further details.

Sub-row assignment in SuiteSparse:GraphBLAS is not as fast as sub-column assignment. If many of the rows of C are to be modified by repeated use of GxB\_Row\_subassign, it can be faster to transpose C first and to use GxB\_Matrix\_subassign or GxB\_Col\_subassign instead.

### 5.8.5 GxB\_Vector\_subassign\_<type>: assign a scalar to a subvector

```
GrB_Info GxB_subassign
                                        // w(I) < mask > = accum (w(I),x)
                                    // input/output vector for results
   GrB_Vector w,
    const GrB_Vector mask,
                                    // optional mask for w(I), unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w(I),x)
    const <type> x,
                                    // scalar to assign to w(I)
                                    // row indices
    const GrB_Index *I,
    const GrB_Index ni,
                                    // number of row indices
    const GrB_Descriptor desc
                                    // descriptor for w(I) and mask
);
```

GxB\_Vector\_subassign\_<type> assigns a single scalar to an entire subvector of the vector w. The operation is exactly like setting a single entry in an n-by-1 matrix, A(I,0) = x, where the column index for a vector is implicitly j=0. For further details of this function, see GxB\_Matrix\_subassign\_<type> in Section 5.8.6.

Unlike GrB\_Vector\_assign\_<type> (see Section 5.9.5), results are not defined if I contains duplicate indices.

#### 5.8.6 GxB\_Matrix\_subassign\_<type>: assign a scalar to a submatrix

```
GrB_Info GxB_subassign
                                        // C(I,J) < Mask > = accum (C(I,J),x)
    GrB_Matrix C,
                                     // input/output matrix for results
    const GrB_Matrix Mask,
                                     // optional mask for C(I,J), unused if NULL
    const GrB_BinaryOp accum,
                                     // optional accum for Z=accum(C(I,J),x)
                                     // scalar to assign to C(I,J)
    const <type> x,
    const GrB_Index *I,
                                     // row indices
    const GrB_Index ni,
                                     // number of row indices
    const GrB_Index *J,
                                     // column indices
    const GrB_Index nj,
                                     // number of column indices
    const GrB_Descriptor desc
                                     // descriptor for C(I,J) and Mask
);
```

GxB\_Matrix\_subassign\_<type> assigns a single scalar to an entire submatrix of C, like the scalar expansion C(I, J)=x in MATLAB. The scalar x is implicitly expanded into a matrix A of size ni by nj, and then the matrix A is assigned to C(I, J) using the same method as in GxB\_Matrix\_subassign. Refer to that function in Section 5.8.2 for further details.

For the accumulation step, the scalar x is typecasted directly into the type of C when the accum operator is not applied to it, or into the ytype of the accum operator, if accum is not NULL, for entries that are already present in C.

The <type> x notation is otherwise the same as GrB\_Matrix\_setElement (see Section 4.8.9). Any value can be passed to this function and its type will be detected, via the \_Generic feature of ANSI C11. For a user-defined type, x is a void \* pointer that points to a memory space holding a single entry of a scalar that has exactly the same user-defined type as the matrix C. This user-defined type must exactly match the user-defined type of C since no typecasting is done between user-defined types.

If a void \* pointer is passed in and the type of the underlying scalar does not exactly match the user-defined type of C, then results are undefined. No error status will be returned since GraphBLAS has no way of catching this error.

Unlike GrB\_Matrix\_assign\_<type> (see Section 5.9.5), results are not defined if I or J contain duplicate indices.

## 5.9 GrB\_assign: submatrix assignment

The methods described in this section are all variations of the form C(I,J)=A, which modifies a submatrix of the matrix C. All methods can be used in their generic form with the single name GrB\_assign. These methods are very similar to their GxB\_subassign counterparts in Section 5.8. They differ primarily in the size of the Mask, and how the GrB\_REPLACE option works. Refer to Section 5.10 for a complete comparison of GxB\_subassign and GrB\_assign.

### 5.9.1 GrB\_Vector\_assign: assign to a subvector

```
GrB_Info GrB_assign
                                    // w<mask>(I) = accum (w(I),u)
                                    // input/output matrix for results
    GrB_Vector w,
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w(I),t)
    const GrB_Vector u,
                                    // first input: vector u
    const GrB_Index *I,
                                    // row indices
    const GrB_Index ni,
                                    // number of row indices
    const GrB_Descriptor desc
                                    // descriptor for w and mask
);
```

GrB\_Vector\_assign operates on a subvector w(I) of w, modifying it with the vector u. The mask vector has the same size as w. The method is identical to GrB\_Matrix\_assign described in Section 5.9.2, where all matrices have a single column each. The only other difference is that the input u in this method is not transposed via the GrB\_INPO descriptor.

#### 5.9.2 GrB\_Matrix\_assign: assign to a submatrix

```
GrB_Info GrB_assign
                                    // C<Mask>(I,J) = accum (C(I,J),A)
    GrB_Matrix C,
                                    // input/output matrix for results
    const GrB_Matrix Mask,
                                    // optional mask for C, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for Z=accum(C(I,J),T)
    const GrB_Matrix A,
                                    // first input: matrix A
    const GrB_Index *I,
                                    // row indices
                                    // number of row indices
    const GrB_Index ni,
    const GrB_Index *J,
                                    // column indices
    const GrB_Index nj,
                                    // number of column indices
    const GrB_Descriptor desc
                                    // descriptor for C, Mask, and A
);
```

GrB\_Matrix\_assign operates on a submatrix S of C, modifying it with the matrix A. It may also modify all of C, depending on the input descriptor desc and the Mask.

Step	GraphBLAS	description
	notation	
1	$\mathbf{S} = \mathbf{C}(\mathbf{I}, \mathbf{J})$	extract $\mathbf{C}(\mathbf{I}, \mathbf{J})$ submatrix
2	$\mathbf{S} = \mathbf{S} \odot \mathbf{A}$	apply the accumulator (but not the mask) to ${f S}$
3	$\mathbf{Z} = \mathbf{C}$	make a copy of $\mathbf{C}$
4	$\mathbf{Z}(\mathbf{I}, \mathbf{J}) = \mathbf{S}$	put the submatrix into $\mathbf{Z}(\mathbf{I}, \mathbf{J})$
5	$\mathbf{C}\langle \mathbf{M} \rangle = \mathbf{Z}$	apply the mask/replace phase to all of ${\bf C}$

In contrast to GxB\_subassign, the Mask has the same as C.

Step 1 extracts the submatrix and then Step 2 applies the accumulator (or S = A if accum is NULL). The Mask is not yet applied.

Step 3 makes a copy of the C matrix, and then Step 4 writes the submatrix S into Z. This is the same as Step 3 of GxB\_subassign, except that it operates on a temporary matrix Z.

Finally, Step 5 writes  $\mathbf{Z}$  back into  $\mathbf{C}$  via the Mask, using the Mask/Replace Phase described in Section 2.3. If  $\mathtt{GrB\_REPLACE}$  is enabled, then all of  $\mathbf{C}$  is cleared prior to writing  $\mathbf{Z}$  via the mask. As a result, the  $\mathtt{GrB\_REPLACE}$  option can delete entries outside the  $\mathbf{C}(\mathbf{I},\mathbf{J})$  submatrix.

#### 5.9.3 GrB\_Col\_assign: assign to a sub-column of a matrix

```
// C<mask>(I,j) = accum (C(I,j),u)
GrB_Info GrB_assign
   GrB_Matrix C,
                                    // input/output matrix for results
                                    // optional mask for C(:,j), unused if NULL
    const GrB_Vector mask,
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(C(I,j),t)
   const GrB_Vector u,
                                    // input vector
    const GrB_Index *I,
                                    // row indices
    const GrB_Index ni,
                                    // number of row indices
    const GrB_Index j,
                                    // column index
    const GrB_Descriptor desc
                                    // descriptor for C(:,j) and mask
);
```

GrB\_Col\_assign modifies a single sub-column of a matrix C. It is the same as GrB\_Matrix\_assign where the index vector J[0]=j is a single column index, and where all matrices in GrB\_Matrix\_assign (except C) consist of a single column.

Unlike GrB\_Matrix\_assign, the mask is a vector with the same size as a single column of C.

The input descriptor GrB\_INPO is ignored; the input vector **u** is not transposed. Refer to GrB\_Matrix\_assign for further details.

#### 5.9.4 GrB\_Row\_assign: assign to a sub-row of a matrix

```
GrB_Info GrB_assign
                                     // C < mask' > (i,J) = accum (C(i,J),u')
                                     // input/output matrix for results
    GrB_Matrix C,
    const GrB_Vector mask,
                                     // optional mask for C(i,:), unused if NULL
                                     // optional accum for z=accum(C(i,J),t)
    const GrB_BinaryOp accum,
    const GrB_Vector u,
                                     // input vector
    const GrB_Index i,
                                     // row index
                                     // column indices
    const GrB_Index *J,
    const GrB_Index nj,
                                     // number of column indices
    const GrB_Descriptor desc
                                     // descriptor for C(i,:) and mask
);
```

GxB\_Row\_subassign modifies a single sub-row of a matrix C. It is the same as GxB\_Matrix\_subassign where the index vector I[0]=i is a single row index, and where all matrices in GxB\_Matrix\_subassign (except C) consist of a single row.

Unlike GrB\_Matrix\_assign, the mask is a vector with the same size as a single row of C.

The input descriptor GrB\_INPO is ignored; the input vector u is not transposed. Refer to GxB\_Matrix\_subassign for further details.

Sub-row assignment in SuiteSparse:GraphBLAS is not as fast as sub-column assignment. If many of the rows of C are to be modified by repeated use of GxB\_Row\_subassign, it can be faster to transpose C first and to use GxB\_Matrix\_subassign or GxB\_Col\_subassign instead. See GxB\_Matrix\_subassign for further details.

### 5.9.5 GrB\_Vector\_assign\_<type>: assign a scalar to a subvector

```
GrB_Info GrB_assign
                                    // w<mask>(I) = accum (w(I),x)
    GrB_Vector w,
                                    // input/output vector for results
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w(I),x)
                                    // scalar to assign to w(I)
    const <type> x,
    const GrB_Index *I,
                                    // row indices
    const GrB_Index ni,
                                    // number of row indices
    const GrB_Descriptor desc
                                    // descriptor for w and mask
);
```

GrB\_Vector\_assign\_<type> assigns a single scalar to an entire subvector of the vector w. The operation is exactly like setting a single entry in an n-by-1 matrix, A(I,0) = x, where the column index for a vector is implicitly j=0. The mask vector has the same size as w. For further details of this function, see GrB\_Matrix\_assign\_<type> in the next section.

In contrast to GxB\_Vector\_subassign\_<type>, results are well-defined if I contains duplicate indices. Duplicate indices are simply ignored.

### 5.9.6 GrB\_Matrix\_assign\_<type>: assign a scalar to a submatrix

```
GrB_Info GrB_assign
                                    // C<Mask>(I,J) = accum (C(I,J),x)
(
    GrB_Matrix C,
                                    // input/output matrix for results
                                    // optional mask for C, unused if NULL
    const GrB_Matrix Mask,
    const GrB_BinaryOp accum,
                                    // optional accum for Z=accum(C(I,J),x)
    const <type> x,
                                    // scalar to assign to C(I,J)
    const GrB_Index *I,
                                    // row indices
    const GrB_Index ni,
                                    // number of row indices
                                    // column indices
    const GrB_Index *J,
    const GrB_Index nj,
                                    // number of column indices
    const GrB_Descriptor desc
                                    // descriptor for C and Mask
);
```

GrB\_Matrix\_assign\_<type> assigns a single scalar to an entire submatrix of C, like the scalar expansion C(I,J)=x in MATLAB. The scalar x is implicitly expanded into a matrix A of size ni by nj, and then the matrix A is assigned to C(I,J) using the same method as in GrB\_Matrix\_assign. Refer to that function in Section 5.9.2 for further details.

The Mask has the same size as C.

For the accumulation step, the scalar  $\mathbf{x}$  is typecasted directly into the type of C when the accum operator is not applied to it, or into the ytype of the accum operator, if accum is not NULL, for entries that are already present in C.

The <type> x notation is otherwise the same as GrB\_Matrix\_setElement (see Section 4.8.9). Any value can be passed to this function and its type will be detected, via the \_Generic feature of ANSI C11. For a user-defined type, x is a void \* pointer that points to a memory space holding a single entry of a scalar that has exactly the same user-defined type as the matrix C. This user-defined type must exactly match the user-defined type of C since no typecasting is done between user-defined types.

If a void \* pointer is passed in and the type of the underlying scalar does not exactly match the user-defined type of C, then results are undefined. No error status will be returned since GraphBLAS has no way of catching this error.

In contrast to GxB\_Matrix\_subassign\_<type>, results are well-defined if I or J contain duplicate indices. Duplicate indices are simply ignored.

# 5.10 Comparing GrB\_assign and GxB\_subassign

SPEC: GxB\_subassign is an extension to the spec.

The GxB\_subassign and GrB\_assign operations are very similar, but they differ in three ways:

- 1. The mask in GxB\_subassign has the same dimensions as w(I) for vectors and C(I,J) for matrices. In GrB\_assign, the mask is the same size as w or C, respectively (except for the row/col variants). The two masks are related. If M is the mask for GrB\_assign, then M(I,J) is the mask for GxB\_subassign. If there is no mask, or if I and J are both GrB\_ALL, then the two masks are the same.
  - For GrB\_Row\_assign and GrB\_Col\_assign, the mask vector is the same size as a row or column of C, respectively. For the corresponding GxB\_Row\_subassign and GxB\_Col\_subassign operations, the mask is the same size as the sub-row C(i, j) or subcolumn C(I, j), respectively.
- 2. They differ in how C is affected in areas outside the C(I,J) submatrix. In GxB\_subassign, the C(I,J) submatrix is the only part of C that can be modified, and no part of C outside the submatrix is ever modified. In GrB\_assign, it is possible to delete entries in C outside the submatrix, but only in one specific manner. Suppose the mask M is present (or, suppose it is not present but GrB\_SCMP is true). After (optionally) complementing the mask, the value of M(i,j) can be 0 for some entry outside the C(I,J) submatrix. If the GrB\_REPLACE descriptor is also true, then GrB\_assign deletes this entry.
- 3. They differ in how duplicate indices are treated in I and J. For both assign and subassign, results are not defined for GrB\_Matrix\_\*assign, GrB\_Vector\_\*assign, GrB\_Row\_\*assign, and GrB\_Col\_\*assign when duplicate indices appear in I or J. The scalar expansion operations, GrB\_\*\_assign\_<type>, are well-defined if duplicate indices appear (the results are the same as if duplicates are removed first from I and J). However, the scalar expansion operations GxB\_\*\_subassign\_<type> are not well-defined if duplicate indices appear in I or J.

GxB\_subassign and GrB\_assign are identical if GrB\_REPLACE is set to its default value of false, and if the masks happen to be the same. The two masks can be the same in two cases: either the Mask input is NULL (and it is not complemented via GrB\_SCMP), or I and J are both GrB\_ALL. In this case, the two algorithms are identical and have the same performance.

GxB\_subassign is much faster than GrB\_assign, when the latter must examine the entire matrix C to delete entries (when GrB\_REPLACE is true), and if it must deal with a much larger Mask matrix. However, both methods have specific uses.

Consider using C(I,J)+=F for many submatrices F (for example, when assembling a finite-element matrix). If the Mask is meant as a specification for which entries of C should appear in the final result, then use GrB\_assign.

If instead the Mask is meant to control which entries of the submatrix C(I,J) are modified by the finite-element F, then use GxB\_subassign. This is particularly useful is the Mask is a template that follows along with the finite-element F, independent of where it is applied to C. Using GrB\_assign would be very difficult in this case since a new Mask, the same size as C, would need to be constructed for each finite-element F.

In GraphBLAS notation, the two methods can be described as follows:

$$\begin{array}{ll} \text{matrix and vector subassign} & \mathbf{C}(\mathbf{I},\mathbf{J})\langle\mathbf{M}\rangle = \mathbf{C}(\mathbf{I},\mathbf{J})\odot\mathbf{A} \\ \text{matrix and vector assign} & \mathbf{C}\langle\mathbf{M}\rangle(\mathbf{I},\mathbf{J}) = \mathbf{C}(\mathbf{I},\mathbf{J})\odot\mathbf{A} \\ \end{array}$$

This notation does not include the details of the GrB\_SCMP and GrB\_REPLACE descriptors, but it does illustrate the difference in the Mask. In the subassign, Mask is the same size as C(I,J) and A. If I[0]=i and J[0]=j, Then Mask(0,0) controls how C(i,j) is modified by the subassign, from the value A(0,0). In the assign, Mask is the same size as C, and Mask(i,j) controls how C(i,j) is modified.

The GxB\_subassign and GrB\_assign functions have the same signatures; they differ only in how they consider the Mask and the GrB\_REPLACE descriptor, and in how duplicate indices are treated for scalar expansion.

Details of each step of the two operations are listed below:

Step	GrB_Matrix_assign	GxB_Matrix_subassign
1	$\mathbf{S} = \mathbf{C}(\mathbf{I}, \mathbf{J})$	$\mathbf{S} = \mathbf{C}(\mathbf{I}, \mathbf{J})$
2	$\mathbf{S} = \mathbf{S} \odot \mathbf{A}$	$\mathbf{S}\langle \mathbf{M} \rangle = \mathbf{S} \odot \mathbf{A}$
3	$\mathbf{Z} = \mathbf{C}$	$\mathbf{C}(\mathbf{I},\mathbf{J})=\mathbf{S}$
4	$\mathbf{Z}(\mathbf{I},\mathbf{J}) = \mathbf{S}$	
5	$\mathbf{C}\langle \mathbf{M}  angle = \mathbf{Z}$	

The Accumulator Phase ( $S \odot A$  in Step 2), described in Section 2.3, is the same in both operations. The result is simply A if accum is NULL. It only applies to the submatrix S, not the whole matrix.

The Mask/Replace Phase, described in Section 2.3 is different:

- For GrB\_Matrix\_assign (Step 5), the mask is applied to all of C. The mask has the same size as C. Just prior to making the assignment via the mask, the GrB\_REPLACE option can be used to clear all of C first. This is the only way in which entries in C that are outside the C(I, J) submatrix can be modified by this operation.
- For  $GxB\_Matrix\_subassign$  (Step 2), the mask is applied to just S. The mask has the same size as C(I,J), S, and A. Just prior to making the assignment via the mask, the  $GrB\_REPLACE$  option can be used to clear S first. No entries in C that are outside the C(I,J) can be modified by this operation. Thus,  $GrB\_REPLACE$  has no effect on entries in C outside the C(I,J) submatrix.

The differences between  $GrB_Matrix_assign$  and  $GxB_Matrix_subassign$  can be seen in Tables 1 and 2. The first table considers the case when the entry  $c_{ij}$  is in the C(I, J) submatrix, and it describes what is computed for both  $GrB_Matrix_assign$  and  $GxB_Matrix_subassign$ . They perform the exact same computation; the only difference is how the value of the mask is specified.

The first column of the table is yes if  $GrB_REPLACE$  is enabled, and a dash otherwise. The second column is yes if an accumulator operator is given, and a dash otherwise. The third column is  $c_{ij}$  if the entry is present in  $\mathbf{C}$ , and a dash otherwise. The fourth column is  $a_{i'j'}$  if the corresponding entry is present in  $\mathbf{A}$ , where  $i = \mathbf{I}(i')$  and  $j = \mathbf{J}(i')$ .

The mask column is 1 if the mask allows C to be modified, and 0 otherwise. This is  $m_{ij}$  for  $GrB\_assign$ , and  $m_{i'j'}$  for  $GxB\_subassign$ , to reflect the difference in the mask, but this difference is not reflected in the table. The value 1 or 0 is the value of the entry in the mask after it is optionally complemented via the  $GrB\_SCMP$  option.

Finally, the last column is the action taken in this case. It is left blank if no action is taken, in which case  $c_{ij}$  is not modified if present, or not inserted into  $\mathbf{C}$  if not present.

repl	accum	$\mathbf{C}$	$\mathbf{A}$	mask	action taken by GrB_assign and GxB_subassign
-	-	$c_{ij}$	$a_{i'j'}$	1	$c_{ij} = a_{i'j'}$ , update
-	-	-	$a_{i'j'}$	1	$c_{ij} = a_{i'j'}$ , insert
-	-	$c_{ij}$	-	1	delete $c_{ij}$ because $a_{i'j'}$ not present
-	-	-	-	1	
-	-	$c_{ij}$	$a_{i'j'}$	0	
-	-	-	$a_{i'j'}$	0	
-	-	$c_{ij}$	-	0	
-	-	-	-	0	
yes	-	$c_{ij}$	$a_{i'j'}$	1	$c_{ij} = a_{i'j'}$ , update
yes	-	-	$a_{i'j'}$	1	$c_{ij} = a_{i'j'}$ , insert
yes	-	$c_{ij}$	-	1	delete $c_{ij}$ because $a_{i'j'}$ not present
yes	-	-	-	1	
yes	-	$c_{ij}$	$a_{i'j'}$	0	delete $c_{ij}$ (because of GrB_REPLACE)
yes	-	-	$a_{i'j'}$	0	
yes	-	$c_{ij}$	-	0	delete $c_{ij}$ (because of GrB_REPLACE)
yes	-	-	-	0	
-	yes	$c_{ij}$	$a_{i'j'}$	1	$c_{ij} = c_{ij} \odot a_{i'j'}$ , apply accumulator
-	yes	-	$a_{i'j'}$	1	$c_{ij} = a_{i'j'}$ , insert
-	yes	$c_{ij}$	-	1	
-	yes	-	-	1	
_	yes	$c_{ij}$	$a_{i'j'}$	0	
-	yes	-	$a_{i'j'}$	0	
-	yes	$c_{ij}$	-	0	
-	yes	-	-	0	
yes	yes	$c_{ij}$	$a_{i'j'}$	1	$c_{ij} = c_{ij} \odot a_{i'j'}$ , apply accumulator
yes	yes	-	$a_{i'j'}$	1	$c_{ij} = a_{i'j'}$ , insert
yes	yes	$c_{ij}$	-	1	
yes	yes	-	-	1	
yes	yes	$c_{ij}$	$a_{i'j'}$	0	delete $c_{ij}$ (because of Grb_REPLACE)
yes	yes	-	$a_{i'j'}$	0	
yes	yes	$c_{ij}$	-	0	delete $c_{ij}$ (because of Grb_REPLACE)
yes	yes	-	_	0	

Table 1: Results of assign and subassign for entries in the  $\mathbf{C}(\mathbf{I},\mathbf{J})$  submatrix

repl	accum	$\mathbf{C}$	$\mathbf{C} = \mathbf{Z}$	mask	action taken by GrB_assign
-	-	$c_{ij}$	$c_{ij}$	1	
-	-	-	-	1	
-	-	$c_{ij}$	$c_{ij}$	0	
-	-	-	-	0	
yes	-	$c_{ij}$	$c_{ij}$	1	
yes	-	-	-	1	
yes	-	$c_{ij}$	$c_{ij}$	0	delete $c_{ij}$ (because of $GrB_REPLACE$ )
yes	-	-	-	0	
-	yes	$c_{ij}$	$c_{ij}$	1	
-	yes	-	-	1	
-	yes	$c_{ij}$	$c_{ij}$	0	
-	yes	-	-	0	
yes	yes	$c_{ij}$	$c_{ij}$	1	
yes	yes	-	-	1	
yes	yes	$c_{ij}$	$c_{ij}$	0	delete $c_{ij}$ (because of $GrB_REPLACE$ )
yes	yes	-	-	0	

Table 2: Results of assign for entries outside the C(I, J) submatrix. Subassign has no effect on these entries.

Table 2 illustrates how  $GrB_assign$  and  $GxB_subassign$  differ for entries outside the submatrix.  $GxB_subassign$  never modifies any entry outside the C(I, J) submatrix, but  $GrB_assign$  can modify them in two cases listed in Table 2. When the  $GrB_REPLACE$  option is selected, and when the Mask(i,j) for an entry  $c_{ij}$  is false (or if the Mask(i,j) is true and  $GrB_sCMP$  is enabled via the descriptor), then the entry is deleted by  $GrB_assign$ .

The fourth column of Table 2 differs from Table 1, since entries in **A** never affect these entries. Instead, for all index pairs outside the  $I \times J$  submatrix, **C** and **Z** are identical (see Step 3 above). As a result, each section of the table includes just two cases: either  $c_{ij}$  is present, or not. This in contrast to Table 1, where each section must consider four different cases.

The GrB\_Row\_assign and GrB\_Col\_assign operations are slightly different. They only affect a single row or column of C. For GrB\_Row\_assign, Table 2 only applies to entries in the single row C(i, J) that are outside the list of indices, J. For GrB\_Col\_assign, Table 2 only applies to entries in the single column C(I,j) that are outside the list of indices, I.

### **5.10.1** Example

The difference between GxB\_subassign and GrB\_assign is illustrated in the following example. Consider the 2-by-2 matrix C where all entries are present.

$$\mathbf{C} = \left[ \begin{array}{cc} 11 & 12 \\ 21 & 22 \end{array} \right]$$

Suppose Grb\_REPLACE is true, and Grb\_SCMP is false. Let the Mask be:

$$\mathbf{M} = \left[ \begin{array}{cc} 1 & 1 \\ 0 & 1 \end{array} \right].$$

Let  $\mathbf{A} = 100$ , and let the index sets be  $\mathbf{I} = 0$  and  $\mathbf{J} = 1$ . Consider the computation  $\mathbf{C}\langle\mathbf{M}\rangle(0,1) = \mathbf{C}(0,1) + \mathbf{A}$ , using the GrB\_assign operation. The result is:

$$\mathbf{C} = \left[ \begin{array}{cc} 11 & 112 \\ - & 22 \end{array} \right].$$

The (0,1) entry is updated and the (1,0) entry is deleted because its Mask is zero. The other two entries are not modified since  $\mathbf{Z} = \mathbf{C}$  outside the submatrix, and those two values are written back into  $\mathbf{C}$  because their Mask values are 1. The (1,0) entry is deleted because the entry  $\mathbf{Z}(1,0) = 21$  is prevented from being written back into  $\mathbf{C}$  since Mask(1,0)=0.

Now consider the analogous  $GxB\_subassign$  operation. The Mask has the same size as A, namely:

$$\mathbf{M} = [1].$$

After computing  $\mathbf{C}(0,1)\langle \mathbf{M} \rangle = \mathbf{C}(0,1) + \mathbf{A}$ , the result is

$$\mathbf{C} = \left[ \begin{array}{cc} 11 & 112 \\ 21 & 22 \end{array} \right].$$

Only the  $\mathbf{C}(\mathbf{I}, \mathbf{J})$  submatrix, the single entry  $\mathbf{C}(0, 1)$ , is modified by  $\mathtt{GxB\_subassign}$ . The entry  $\mathbf{C}(1, 0) = 21$  is unaffected by  $\mathtt{GxB\_subassign}$ , but it is deleted by  $\mathtt{GrB\_assign}$ .

### 5.10.2 Performance of GxB\_subassign, GrB\_assign and GrB\_\*\_setElement

When SuiteSparse:GraphBLAS uses non-blocking mode, the modifications to a matrix by GxB\_subassign, GrB\_assign, and GrB\_\*\_setElement can postponed, and computed all at once later on. This has a huge impact on performance.

A sequence of assignments is fast if their completion can be postponed for as long as possible, or if they do not modify the pattern at all. Modifying the pattern can be costly, but it is fast if non-blocking mode can be fully exploited.

Consider a sequence of t submatrix assignments C(I,J)=C(I,J)+A to an n-by-n matrix C where each submatrix A has size a-by-a with s entries, and where C starts with k entries.

If blocking mode is enabled, or if the sequence requires the matrix to be completed after each assignment, each of the t assignments takes  $O(a + s \log n)$  time to process the A matrix and then  $O(n + k + s \log s)$  time to complete C. The latter step uses  $\texttt{GrB}_*=\texttt{build}$  to build an update matrix and then merge it with C. This step does not occur if the sequence of assignments does not add new entries to the pattern of C, however. Assuming in the worst case that the pattern does change, the total time is  $O(t [a + s \log n + n + k + s \log s])$ .

If the sequence can be computed with all updates postponed until the end of the sequence, then the total time is no worse than  $O(a+s\log n)$  to process each A matrix, for t assignments, and then a single build at the end, taking  $O(n+k+st\log st)$  time. The total time is  $O(t\left[a+s\log n\right]+(n+k+st\log st))$ . If no new entries appear in C the time drops to  $O(t\left[a+s\log n\right])$ , and in this case, the time for both methods is the same; both are equally efficient.

A few simplifying assumptions are useful to compare these times. Nearly all graphs of n nodes that arise in practice have O(n) edges, and most graphs have a constant bound on the degree of each node. The asymptotic bounds assume a worst-case scenario where C has a least some dense columns (thus the  $\log n$  terms). If these are not present, if both t and k are O(n), and if a and s are constants, then the total time with blocking mode becomes  $O(n^2)$ , assuming the pattern of C changes at each assignment. This very high for a sparse graph problem. In contrast, the non-blocking time becomes  $O(n \log n)$  under these same assumptions, which is asymptotically much faster.

The difference in practice can be very dramatic, since n can be many millions for sparse graphs that can be handled on a commodity laptop.

The following guidelines should be considered when using GxB\_subassign, GrB\_assign and GrB\_\*\_setElement.

- 1. A sequence of assignments that does not modify the pattern at all is fast, taking as little as  $\Omega(1)$  time per entry modified. The worst case time complexity is  $O(\log n)$  per entry, assuming they all modify a dense column of C with n entries, which can occur in practice. It is more common, however, that most columns of C have a constant number of entries, independent of n. No work is ever left pending when the pattern of C does not change.
- 2. A sequence of assignments that modifies the entries that already exist in the pattern of a matrix, or adds new entries to the pattern (using the same accum operator), but does not delete any entries, is fast. The matrix is not completed until the end of the sequence.
- 3. Similarly, a sequence that modifies existing entries, or deletes them, but does not add new ones, is also fast. This sequence can also repeatedly delete pre-existing entries and then reinstate them and still be fast. The matrix is not completed until the end of the sequence.
- 4. A sequence that mixes assignments of types (2) and (3) above can be costly, since the matrix may need to be completed after each assignment. The time complexity can become quadratic in the worst case.
- 5. However, any single assignment takes no more than  $O(a+s\log n+n+k+s\log s)$  time, even including the time for a matrix completion, where C is n-by-n with k entries and A is a-by-a with s entries. This time is essentially linear in the size of the matrix C, if A is relatively small and sparse compared with C. In this case, n+k are the two dominant terms.
- 6. In general, GxB\_subassign is faster than GrB\_assign. If GrB\_REPLACE is used with GrB\_assign, the entire matrix C must be traversed. This is much slower than GxB\_subassign, which only needs to examine the C(I, J) submatrix. Furthermore, GrB\_assign must deal with a much larger Mask matrix, whereas GxB\_subassign has a smaller mask. Since its mask is smaller, GxB\_subassign takes less time than GrB\_assign to access the mask.

Submatrix assignment in SuiteSparse:GraphBLAS is extremely efficient, even without considering the advantages of non-blocking mode discussed in Section 5.10. Consider assigning a large submatrix C(I,J)=A where C is the Freescale2 matrix from the SuiteSparse Collection [DH11], of size 3 million by 3 million, with 14.3 million nonzeros. With the vectors I=randperm(n,5500) and J=randperm(n,7000) and A a random sparse matrix with 38,500 nonzeros, C(I,J)=A takes 87 seconds in MATLAB. The same computation takes 0.74 seconds in SuiteSparse:GraphBLAS, a speedup of over 100. This is after finishing all pending computations in GraphBLAS and returning result to MATLAB as a valid MATLAB sparse matrix. The dominant time complexity for GraphBLAS is O(n+k), where n is the dimension of C and C is its number of nonzeros. As a comparison, MATLAB takes just 0.42 seconds to compute C+C for this matrix, which also takes time linear in the size of the matrix data structure, O(n+k).

 $<sup>^1\</sup>mathrm{All}$  performance measurements in this document were done on a MacBook Pro, 2.8 GHz Intel Core i7, 16 GB Ram, OSX 10.11.6, clang 8.0.0, MATLAB R2017A.

## 5.11 GrB\_apply: apply a unary operator

The GrB\_apply function is the generic name for two specific functions: GrB\_Vector\_apply and GrB\_Matrix\_apply. The generic name appears in the function prototypes, but the specific function name is used when describing each variation. When discussing features that apply to both versions, the simple name GrB\_apply is used.

### 5.11.1 GrB\_Vector\_apply: apply a unary operator to a vector

GrB\_Vector\_apply applies a unary operator to the entries of a vector, analogous to  $\mathbf{t} = op(\mathbf{u})$  in MATLAB except the operator op is only applied to entries in the pattern of  $\mathbf{u}$ . Implicit values outside the pattern of  $\mathbf{u}$  are not affected. The entries in  $\mathbf{u}$  are typecasted into the xtype of the unary operator. The vector  $\mathbf{t}$  has the same type as the ztype of the unary operator. The final step is  $\mathbf{w}\langle \mathbf{m} \rangle = \mathbf{w} \odot \mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices.

#### 5.11.2 GrB\_Matrix\_apply: apply a unary operator to a matrix

GrB\_Matrix\_apply applies a unary operator to the entries of a matrix, analogous to T = op(A) in MATLAB except the operator op is only applied to entries in the pattern of A. Implicit values outside the pattern of A are not affected. The input matrix A may be transposed first. The entries in A are typecasted into the xtype of the unary operator. The matrix T has the same type as the ztype of the unary operator. The final step is  $C\langle M \rangle = C \odot T$ , as described in Section 2.3.

The built-in  $GrB_IDENTITY_T$  operators (one for each built-in type T) are very useful when combined with this function, enabling it to compute  $C\langle M \rangle = C \odot A$ . This makes  $GrB_apply$  a direct interface to the accumulator/mask function for both matrices and vectors.

In SuiteSparse:GraphBLAS, this method is particularly efficient with built-in types. If the type of C and A are the same, and if A is not transposed via the descriptor, then T is a pure shallow copy of A, taking only O(1) time and memory. The output matrix C is never a shallow copy of T or A.

To compute  $\mathbf{C}\langle\mathbf{M}\rangle=\mathbf{A}$  or  $\mathbf{C}\langle\mathbf{M}\rangle=\mathbf{C}\odot\mathbf{A}$  for user-defined types, the user application would need to define an identity operator for the type. Since GraphBLAS cannot detect that it is an identity operator, it must call the operator to make the full copy T=A and apply the operator to each entry of the matrix or vector.

The other GraphBLAS operation that provides a direct interface to the accumulator/mask function is GrB\_transpose, which does not require an operator to perform this task. As a result, GrB\_transpose can be used as an efficient and direct interface to the accumulator/mask function for both built-in and user-defined types. However, it is only available for matrices, not vectors.

## 5.12 GxB\_select: apply a select operator

The GxB\_select function is the generic name for two specific functions: GxB\_Vector\_select and GxB\_Matrix\_select. The generic name appears in the function prototypes, but the specific function name is used when describing each variation. When discussing features that apply to both versions, the simple name GxB\_select is used.

**SPEC:** The GxB\_select operation and GxB\_SelectOp operator are extensions to the spec.

### 5.12.1 GxB\_Vector\_select: apply a select operator to a vector

```
GrB_Info GxB_select
                                    // w<mask> = accum (w, op(u,k))
                                    // input/output vector for results
    GrB_Vector w,
    const GrB_Vector mask,
                                    // optional mask for w, unused if NULL
    const GrB_BinaryOp accum,
                                    // optional accum for z=accum(w,t)
                                    // operator to apply to the entries
    const GxB_SelectOp op,
    const GrB_Vector u,
                                    // first input: vector u
    const void *k,
                                    // optional input for the select operator
    const GrB_Descriptor desc
                                    // descriptor for w and mask
);
```

GxB\_Vector\_select applies a select operator to the entries of a vector, analogous to t = u.\*op(u) in MATLAB except the operator op is only applied to entries in the pattern of u. Implicit values outside the pattern of u are not affected. If the operator is not type-generic, the entries in u are type-casted into the xtype of the select operator. The vector t has the same type and size as u. The final step is  $\mathbf{w}\langle \mathbf{m}\rangle = \mathbf{w}\odot \mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices.

This operation operates on vectors just as if they were m-by-1 matrices, except that GraphBLAS never transposes a vector via the descriptor. The op is passed n=1 as the number of columns. Refer to the next section on GxB\_Matrix\_select for more details.

#### 5.12.2 GxB\_Matrix\_select: apply a select operator to a matrix

```
// C<Mask> = accum (C, op(A,k)) or op(A',k)
GrB_Info GxB_select
    GrB_Matrix C,
                                    // input/output matrix for results
                                    // optional mask for C, unused if NULL
    const GrB_Matrix Mask,
    const GrB_BinaryOp accum,
                                    // optional accum for Z=accum(C,T)
                                    // operator to apply to the entries
    const GxB_SelectOp op,
    const GrB_Matrix A,
                                    // first input: matrix A
                                    // optional input for the select operator
    const void *k,
    const GrB_Descriptor desc
                                    // descriptor for C, mask, and A
);
```

GxB\_Matrix\_select applies a select operator to the entries of a matrix, analogous to T = A .\* op(A) in MATLAB except the operator op is only applied to entries in the pattern of A. Implicit values outside the pattern of A are not affected. The input matrix A may be transposed first. If the operator is not type-generic, the entries in A are typecasted into the xtype of the select operator. The final step is  $C\langle M \rangle = C \odot T$ , as described in Section 2.3.

The matrix T has the same size and type as A (or the transpose of A if the input is transposed via the descriptor). The entries of T are a subset of those of A. Each entry A(i,j) of A is passed to the op, as  $z = f(i,j,m,n,a_{ij},k)$ , where A is m-by-n. If A is transposed first then the operator is applied to entries in the transposed matrix, A'. If z is returned as true, then the entry is copied into T, unchanged. If it returns false, the entry does not appear in T.

For user-defined select operators, the argument k is passed to the operator unchanged. For built-in operators, k is a pointer to an int64\_t scalar that refers to the kth diagonal of the matrix. The value k=0 specifies the main diagonal of the matrix, k=1 is the +1 diagonal (the entries just above the main diagonal), k=-1 is the -1 diagonal, and so on. Note that k must be passed as a pointer to int64\_t, not merely as an integer. The parameter k is not used by GxB\_NONZERO and may be passed as GrB\_NULL.

The action of GxB\_select with the built-in select operators is described in the table below. The MATLAB analogs are precise for tril and triu, but shorthand for the other operations. The MATLAB diag function returns a column with the diagonal, if A is a matrix, whereas the matrix T in GxB\_select always has same size as A (or its transpose if the GrB\_INPO is set to GrB\_TRAN). In the MATLAB analog column, diag is as if it operates like GxB\_select, where T is a matrix.

GraphBLAS	MATLAB	
name	analog	
GxB_TRIL	T=tril(A,k)	Entries in T are the entries on and below the
		kth diagonal of A.
GxB_TRIU	T=triu(A,k)	Entries in T are the entries on and above the
		kth diagonal of A.
GxB_DIAG	T=diag(A,k)	Entries in T are the entries on the kth diagonal
		of A.
GxB_OFFDIAG	T=A-diag(A,k)	Entries in T are all entries not on the kth di-
		agonal of A.
GxB_NONZERO	$T=A(A^{\sim}=0)$	Entries in T are all entries in A that have
		nonzero value.

#### 5.13 GrB reduce: reduce to a vector or scalar

The generic function name <code>GrB\_reduce</code> may be used for all specific functions discussed in this section. When the details of a specific function are discussed, the specific name is used for clarity.

#### 5.13.1 GrB\_Matrix\_reduce\_<op>: reduce a matrix to a vector

GrB\_Matrix\_reduce\_<op> is a generic name for two specific methods. Both methods reduce a matrix to a column vector using an operator, roughly analogous to t = sum (A') in MATLAB, in the default case, where t is a column vector. By default, the method reduces across the rows to obtain a column vector; use GrB\_TRAN to reduce down the columns.

GrB\_Matrix\_reduce\_BinaryOp relies on a binary operator for the reduction: the fourth argument reduce, a GrB\_BinaryOp. All three domains of the operator must be the same. GrB\_Matrix\_reduce\_Monoid performs the same reduction using a GrB\_Monoid as its fourth argument. In both cases the reduction operator must be commutative and associative. Otherwise the results are undefined.

The input matrix A may be transposed first. Its entries are then typecast into the type of the reduce operator or monoid. The reduction is applied to all entries in A (i,:) to produce the scalar t (i). This is done without the use of the identity value of the monoid. If the ith row A (i,:) has no entries, then (i) is not an entry in t and its value is implicit. If A (i,:) has a single entry, then that is the result t (i) and reduce is not applied at all for the ith row. Otherwise, multiple entries in row A (i,:) are reduced via the reduce operator or monoid to obtain a single scalar, the result t (i).

The final step is  $\mathbf{w}\langle \mathbf{m} \rangle = \mathbf{w} \odot \mathbf{t}$ , as described in Section 2.3, except that all the terms are column vectors instead of matrices.

#### 5.13.2 GrB\_Vector\_reduce\_<type>: reduce a vector to a scalar

GrB\_Vector\_reduce\_<type> reduces a vector to a scalar, analogous to t = sum (u) in MATLAB, except that in GraphBLAS any commutative and associative monoid can be used in the reduction.

The reduction operator is a commutative and associative monoid with an identity value. Results are undefined if the monoid does not have these properties. This function differs from <code>GrB\_Matrix\_reduce\_BinaryOp</code> (which reduces a matrix to a vector) in that it requires a valid monoid additive identity value. If the vector u has no entries, that identity value is copied into the scalar t. Otherwise, all of the entries in the vector are reduced to a single scalar using the <code>reduce</code> operator.

The scalar type is any of the built-in types, or a user-defined type. In the function signature it is a C type: bool, int8\_t, ... float, double, or void \* for a user-defined type. The user-defined type must be identical to the type of the vector u. This cannot be checked by GraphBLAS and thus results are undefined if the types are not the same.

The descriptor is unused, but it appears in case it is needed in future versions of the GraphBLAS API Specification. This function has no mask so its accumulator/mask step differs from the other GraphBLAS operations. It does not use the methods described in Section 2.3, but uses the following method instead.

If accum is NULL, then the scalar t is typecast into the type of c, and c = t is the final result. Otherwise, the scalar t is typecast into the ytype of the accum operator, and the value of c (on input) is typecast into the xtype of the accum operator. Next, the scalar z = accum (c,t) is computed, of the ztype of the accum operator. Finally, z is typecast into the final result, c.

**Forced completion:** All computations for the vector **u** are guaranteed to be finished when the method returns.

#### 5.13.3 GrB\_Matrix\_reduce\_<type>: reduce a matrix to a scalar

GrB\_Matrix\_reduce\_<type> reduces a matrix A to a scalar, roughly analogous to t = sum (A (:)) in MATLAB. This function is identical to reducing a vector to a scalar, since the positions of the entries in a matrix or vector have no effect on the result. Refer to the reduction to scalar described in the previous Section 5.13.2.

**Forced completion:** All computations for the matrix **A** are guaranteed to be finished when the method returns.

## 5.14 GrB\_transpose: transpose a matrix

GrB\_transpose transposes a matrix A, just like the array transpose T = A.' in MATLAB. The internal result matrix T = A' (or merely T = A if A is transposed via the descriptor) has the same type as A. The final step is  $\mathbf{C}\langle\mathbf{M}\rangle = \mathbf{C}\odot\mathbf{T}$ , as described in Section 2.3, which typecasts T as needed and applies the mask and accumulator.

To be consistent with the rest of the GraphBLAS API Specification regarding the descriptor, the input matrix A may be transposed first. It may seem counter-intuitive, but this has the effect of not doing any transpose at all. As a result,  $GrB\_transpose$  is useful for more than just transposing a matrix. It can be used as a direct interface to the accumulator/mask operation,  $C\langle M \rangle = C \odot A$ . This step also does any typecasting needed, so  $GrB\_transpose$  can be used to typecast a matrix A into another matrix C. To do this, simply use NULL for the Mask and accum, and provide a non-default descriptor desc that sets the transpose option:

```
// C = typecasted copy of A
GrB_Descriptor_set (desc, GrB_INPO, GrB_TRAN) ;
GrB_transpose (C, NULL, NULL, A, desc) ;
```

If the types of C and match, then the above two lines of code are the same as GrB\_Matrix\_dup (&C, A), except that for GrB\_transpose the matrix C must already exist and be the right size. If C does not exist, the work of GrB\_Matrix\_dup can be replicated with this:

```
// C = create an exact copy of A, just like GrB_Matrix_dup
GrB_Matrix C;
GrB_Type type;
GrB_Index nrows, ncols;
GrB_Descriptor desc;
GxB_Matrix_type (&type, A);
GrB_Matrix_nrows (&nrows, A);
```

```
GrB_Matrix_ncols (&ncols, A) ;
GrB_Matrix_new (&C, type, nrows, ncols) ;
GrB_Descriptor_new (&desc) ;
GrB_Descriptor_set (desc, GrB_INPO, GrB_TRAN) ;
GrB_transpose (C, NULL, NULL, A, desc) ;
```

Since the input matrix A is transposed by the descriptor, SuiteSparse:Graph-BLAS does the right thing and does not transpose the matrix at all. Since T = A is not typecasted, SuiteSparse:GraphBLAS can construct T internally in O(1) time and using no memory at all. This makes  $Grb_transpose$  a fast and direct interface to the accumulator/mask function in GraphBLAS.

This example is of course overkill, since the work can all be done by a single call to the <code>GrB\_Matrix\_dup</code> function. However, the <code>GrB\_Matrix\_dup</code> function can only create <code>C</code> as an exact copy of <code>A</code>, whereas variants of the code above can do many more things with these two matrices. For example, the <code>type</code> in the example can be replaced with any other type, perhaps selected from another matrix or from an operator.

Consider the following code excerpt, which uses  $GrB\_transpose$  to remove all diagonal entries from a square matrix. It first creates a diagonal Mask, which is complemented so that  $\mathbf{C}\langle \neg \mathbf{M}\rangle = \mathbf{A}$  does not modify the diagonal of  $\mathbf{C}$ . The REPLACE ensures that  $\mathbf{C}$  is cleared first, and then  $\mathbf{C}\langle \neg \mathbf{M}\rangle = \mathbf{A}$  modifies all entries in  $\mathbf{C}$  where the mask  $\mathbf{M}$  is false. These correspond to all the off-diagonal entries. The descriptor ensures that  $\mathbf{A}$  is not transposed at all. The Mask can have any pattern, of course, and wherever it is set true, the corresponding entries in  $\mathbf{A}$  are deleted from the copy  $\mathbf{C}$ .

```
// remove all diagonal entries from the matrix A
// Mask = speye (n);
GrB_Matrix_new (&Mask, GrB_BOOL, n, n);
for (int64_t i = 0; i < n; i++)
{
        GrB_Matrix_setElement (Mask, (bool) true, i, i);
}
// C<~Mask> = A, clearing C first. No transpose.
GrB_Descriptor_new (&desc);
GrB_Descriptor_set (desc, GrB_INPO, GrB_TRAN);
GrB_Descriptor_set (desc, GrB_MASK, GrB_SCMP);
GrB_Descriptor_set (desc, GrB_OUTP, GrB_REPLACE);
GrB_transpose (A, Mask, NULL, A, desc);
```

# 6 Examples

Six examples on how to use GraphBLAS are described below: (1) performing a breadth-first search, (2) finding a maximal independent set, (3) creating a random matrix, (4) creating a finite-element matrix, (5) reading a matrix from a file, and (6) complex numbers as a user-defined type. The complete set of programs appears in the Demo directory in SuiteSparse:GraphBLAS.

#### 6.1 Breadth-first search

The bfs examples in the Demo folder provide several examples of how to compute a breadth-first search (BFS) in GraphBLAS. The bfs5m function starts at a given source node s of an undirected graph with n nodes. The graph is represented as a symmetric n-by-n Boolean matrix, A. The matrix A can actually have any type; if it is not Boolean (bool in C, or GrB\_BOOL in GraphBLAS), it is typecasted to Boolean by the semiring, where zero is false and nonzero is true.

The vector  $\mathbf{v}$  of size  $\mathbf{n}$  holds the level of each node in the BFS, where  $\mathbf{v}(\mathtt{i})=0$  if the node has not yet been seen. This particular value makes  $\mathbf{v}$  useful for another role. It can be used as a Boolean mask, since 0 is false and nonzero is true. Initially the entire  $\mathbf{v}$  vector is zero.

The vector **q** is the set of nodes just discovered at the current level, where **q(i)=true** if node **i** is in the current level. It starts out with just a single entry set to true, **q(s)**, the starting node.

Each iteration of the BFS consists of three calls to GraphBLAS. The first one uses q as a mask. It modifies all positions in v where q is true, setting them all to the current level. No accumulator or descriptor are used. Since GrB\_REPLACE is not used and I=GrB\_ALL, GxB\_subassign and GrB\_assign are identical; either can be used in this step:

```
// v < q > = level, using vector assign with q as the mask GrB_assign (v, q, NULL, level, GrB_ALL, n, NULL) ;
```

The next call to GraphBLAS is the heart of the algorithm:

```
// q<!v> = A \mid |.\&\& q ; finds all the unvisited 
// successors from current q, using !v as the mask 
GrB_mxv (q, v, NULL, Boolean, A, q, desc);
```

The vector  $\mathbf{q}$  is all the set of nodes at the current level. Suppose  $\mathbf{q}(\mathbf{j})$  is true, and it has a neighbor  $\mathbf{i}$ . Then  $\mathbf{A}(\mathbf{i},\mathbf{j})=1$ , and the dot product of  $\mathbf{A}(\mathbf{i},\mathbf{i})*\mathbf{q}$  using the OR-AND semiring will use the AND multiplier on these two terms,  $\mathbf{A}(\mathbf{i},\mathbf{j})$  AND  $\mathbf{q}(\mathbf{j})$ , resulting in a value true. The OR monoid will "sum" up all the results in this single row  $\mathbf{i}$ . If the result is a column vector  $\mathbf{t}=\mathbf{A}*\mathbf{q}$ , then this  $\mathbf{t}(\mathbf{i})$  will be true. The vector  $\mathbf{t}$  will be true for any node adjacent to any node in the set  $\mathbf{q}$ .

Some of these neighbors of the nodes in q have already been visited by the BFS, either in the current level or in a prior level. These results must be discarded; what is desired is the set of all nodes i for which t(i) is true, and yet v(i) is still zero.

Enter the mask. The vector  $\mathbf{v}$  is complemented for use a mask, via the desc descriptor. This means that wherever the vector is true, that position in the result is protected and will not be modified by the assignment. Only where  $\mathbf{v}$  is false will the result be modified. This is exactly the desired result, since these represent newly seen nodes for the next level of the BFS. A node  $\mathbf{k}$  already visited will have a nonzero  $\mathbf{v}(\mathbf{k})$ , and thus  $\mathbf{q}(\mathbf{k})$  will not be modified by the assignment.

The result t is written back into the vector q, through the mask, but to do this correctly, another descriptor parameter is used: GrB\_REPLACE. The vector q was used to compute t=A\*q, and after using it to compute t, the entire q vector needs to be cleared. Only new nodes are desired, for the next level. This is exactly what the REPLACE option does.

As a result, the vector **q** now contains the set of nodes at the new level of the BFS. It contains all those nodes (and only those nodes) that are neighbors of the prior set and that have not already been seen in any prior level.

Finally, a single call to GraphBLAS computes the OR for all entries in q, into a single scalar, successor. This value is true if q contains any value true, or false otherwise. If it is false, the BFS can terminate.

```
GrB_reduce (&successor, NULL, Lor, q, NULL) ;
```

The bfs5m function is a modified version from *The GraphBLAS C API Specification* [BMM<sup>+</sup>17]. The method here uses GrB\_mxv instead of GrB\_vxm.

Another method for computing the BFS is in the bfs6 function in the Demo folder. It uses GrB\_apply and a unary operator to set the levels of the newly discovered nodes, instead of GrB\_assign.

```
GrB_Info bfs5m
                          // BFS of a graph (using vector assign & reduce)
   GrB_Vector *v_output,
                          // v [i] is the BFS level of node i in the graph
   const GrB_Matrix A,
                          // input graph, treated as if boolean in semiring
   GrB_Index s
                          // starting node of the BFS
)
{
   GrB_Info info ;
   GrB_Index n ;
                                         // # of nodes in the graph
   GrB_Vector q = NULL ;
                                         // nodes visited at each level
   GrB_Vector v = NULL ;
                                        // result vector
   GrB_Monoid Lor = NULL ;
                                        // Logical-or monoid
                                        // Boolean semiring
   GrB_Semiring Boolean = NULL ;
   GrB_Descriptor desc = NULL ;
                                        // Descriptor for mxv
                                        // n = # of rows of A
   GrB_Matrix_nrows (&n, A) ;
   GrB_Vector_new (&q, GrB_BOOL, n) ; // Vector<bool> q(n) = false
   for (int32_t i = 0 ; i < n ; i++) GrB_Vector_setElement (v, 0, i) ;</pre>
   GrB_Vector\_setElement (q, true, s) ; // q[s] = true, false elsewhere
   GrB_Monoid_new (&Lor, GrB_LOR, (bool) false);
   GrB_Semiring_new (&Boolean, Lor, GrB_LAND) ;
   GrB_Descriptor_new (&desc) ;
   GrB_Descriptor_set (desc, GrB_MASK, GrB_SCMP) ;  // invert the mask
   GrB_Descriptor_set (desc, GrB_OUTP, GrB_REPLACE) ; // clear q first
   bool successor = true ; // true when some successor found
   for (int32_t level = 1 ; successor && level <= n ; level++)</pre>
       // v < q > = level, using vector assign with q as the mask
       GrB_assign (v, q, NULL, level, GrB_ALL, n, NULL) ;
       // q<!v> = A ||.&& q ; finds all the unvisited successors from current
       // q, using !v as the mask
       GrB_mxv (q, v, NULL, Boolean, A, q, desc) ;
       // successor = ||(q)
       GrB_reduce (&successor, NULL, Lor, q, NULL) ;
   v_output = v;
                          // return result
   GrB_free (&q) ;
                          // free workspace
   GrB_free (&Lor) ; GrB_free (&Boolean) ; GrB_free (&desc) ;
   return (GrB_SUCCESS) ;
}
```

## 6.2 Maximal independent set

The maximal independent set problem is to find a set of nodes S such that no two nodes in S are adjacent to each other (an independent set), and all nodes not in S are adjacent to at least one node in S (and thus S is maximal since it cannot be augmented by any node while remaining an independent set). The mis function in the Demo folder solves this problem using Luby's method [Lub86]. The key operations in the method are replicated on the next page.

The gist of the algorithm is this. In each phase, all candidate nodes are given a random score. If a node has a score higher than all its neighbors, then it is added to the independent set. All new nodes added to the set cause their neighbors to be removed from the set of candidates. The process must be repeated for multiple phases until no new nodes can be added. This is because in one phase, a node  $\mathbf i$  might not be added because one of its neighbors  $\mathbf j$  has a higher score, yet that neighbor  $\mathbf j$  might not be added because one of its neighbors  $\mathbf k$  is added to the independent set instead. The node  $\mathbf j$  is no longer a candidate and can never be added to the independent set, but node  $\mathbf i$  could be added to S in a subsequent phase.

The initialization step, before the while loop, computes the degree of each node with a PLUS reduction. The set of candidates is Boolean vector, the ith component is true if node i is a candidate. A node with no neighbors causes the algorithm to stall, so these nodes are not candidates. Instead, they are immediately added to the independent set, represented by another Boolean vector iset. Both steps are done with an assign, using the degree as a mask, except the assignment to iset uses the complement of the mask, via the sr\_desc descriptor. Finally, the GrB\_Vector\_nvals statement counts how many candidates remain.

Each phase of Luby's algorithm consists of nine calls to GraphBLAS operations. Not all of them are described here since they are commented in the code itself. The two matrix-vector multiplications are the important parts and also take the most time. They also make interesting use of semirings and masks. The first one computes the largest score of all the neighbors of each node in the candidate set:

```
// compute the max probability of all neighbors
GrB_mxv (neighbor_max, candidates, NULL, maxSelect2nd, A, prob, r_desc);
```

```
// compute the degree of each node
GrB_reduce (degrees, NULL, NULL, GrB_PLUS_FP64, A, NULL) ;
// singletons are not candidates; they are added to iset first instead
// candidates[degree != 0] = 1
GrB_assign (candidates, degrees, NULL, true, GrB_ALL, n, NULL);
// add all singletons to iset
// iset[degree == 0] = 1
GrB_assign (iset, degrees, NULL, true, GrB_ALL, n, sr_desc) ;
// Iterate while there are candidates to check.
GrB_Index nvals ;
GrB_Vector_nvals (&nvals, candidates);
while (nvals > 0)
    // compute a random probability scaled by inverse of degree
    GrB_apply (prob, candidates, NULL, set_random, degrees, r_desc) ;
    // compute the max probability of all neighbors
    GrB_mxv (neighbor_max, candidates, NULL, maxSelect2nd, A, prob, r_desc)
    // select node if its probability is > than all its active neighbors
    GrB_eWiseAdd (new_members, NULL, NULL, GrB_GT_FP64, prob, neighbor_max,
                                                                    NULL) ;
    // add new members to independent set.
    GrB_eWiseAdd (iset, NULL, NULL, GrB_LOR, iset, new_members, NULL) ;
    // remove new members from set of candidates c = c & !new
    GrB_apply (candidates, new_members, NULL, GrB_IDENTITY_BOOL,
        candidates, sr_desc);
    GrB_Vector_nvals (&nvals, candidates) ;
    if (nvals == 0) { break ; }
                                                 // early exit condition
    // Neighbors of new members can also be removed from candidates
    GrB_mxv (new_neighbors, candidates, NULL, Boolean, A,
        new_members, NULL) ;
    GrB_apply (candidates, new_neighbors, NULL, GrB_IDENTITY_BOOL,
        candidates, sr_desc);
    GrB_Vector_nvals (&nvals, candidates);
}
```

A is a Boolean matrix and prob is a sparse real vector (of type FP32), where prob(j) is nonzero only if node j is a candidate. The maxSelect2nd semiring uses z=SECOND(x,y) as the multiplier operator. The row A(i,:) is the adjacency of node i, and the dot product A(i,:)\*prob applies the SECOND operator on all entries that appear in the intersection of A(i,:) and prob, z=SECOND(A(i,j),prob(j)) which is just prob(j) if A(i,j) is present. If A(i,j) not an explicit entry in the matrix, then this term is not computed and does not take part in the reduction by the MAX monoid.

Thus, each term z=SECOND(A(i,j),prob(j)) is the score, prob(j), of all neighbors j of node i that have a score. Node j does not have a score if it is not also a candidate and so this is skipped. These terms are then "summed" up by taking the maximum score, using MAX as the additive monoid.

Finally, the results of this matrix-vector multiply are written to the result, neighbor\_max. The r\_desc descriptor has the REPLACE option enabled. Since neighbor\_max does not also take part in the computation A\*prob, it is simply cleared first. Next, is it modified only in those positions i where candidates(i) is true, using candidates as a mask. This sets the neighbor\_max only for candidate nodes, and leaves the other components of neighbor\_max as zero (implicit values not in the pattern of the vector).

All of the above work is done in a single matrix-vector multiply, with an elegant use of the maxSelect2nd semiring coupled with a mask. The matrix-vector multiplication is described above as if it uses dot products of rows of A with the column vector prob, but SuiteSparse:GraphBLAS does not compute it that way. Sparse dot products are much slower the optimal method for multiplying a sparse matrix times a sparse vector. The result is the same, however.

The second matrix-vector multiplication is more straight-forward. Once the set of new members in the independent is found, it is used to remove all neighbors of those new members from the set of candidates.

The resulting method is very efficient. For the Freescale2 matrix, the algorithm finds an independent set of size 1.6 million in 1.7 seconds (on the same MacBook Pro referred to in Section 6.1), taking four iterations of the while loop. For comparison, removing its diagonal entries (required for the algorithm to work) takes 0.3 seconds in GraphBLAS (see Section 5.14), and simply transposing the matrix takes 0.24 seconds in both MATLAB and GraphBLAS.

## 6.3 Creating a random matrix

The random\_matrix function in the Demo folder generates a random matrix with a specified dimension and number of entries, either symmetric or unsymmetric, and with or without self-edges (diagonal entries in the matrix). It relies on simple\_rand\* functions in the Demo folder to provide a portable random number generator that creates the same sequence on any computer and operating system.

random\_matrix can use one of two methods: GrB\_Matrix\_setElement and GrB\_Matrix\_build. The former method is very simple to use:

```
GrB_Matrix_new (&A, GrB_FP64, nrows, ncols);
for (int64_t k = 0; k < ntuples; k++)
{
    GrB_Index i = simple_rand_i () % nrows;
    GrB_Index j = simple_rand_i () % ncols;
    if (no_self_edges && (i == j)) continue;
    double x = simple_rand_x ();
    // A (i,j) = x
    GrB_Matrix_setElement (A, x, i, j);
    if (make_symmetric)
    {
        // A (j,i) = x
        GrB_Matrix_setElement (A, x, j, i);
    }
}</pre>
```

The above code can generate a million-by-million sparse double matrix with 200 million entries in 66 seconds (6 seconds of which is the time to generate the random i, j, and x), including the time to finish all pending computations. The user application does not need to create a list of all the tuples, nor does it need to know how many entries will appear in the matrix. It just starts from an empty matrix and adds them one at a time in arbitrary order. GraphBLAS handles the rest. This method is not feasible in MATLAB.

The next method uses GrB\_Matrix\_build. It is more complex to use than setElement since it requires the user application to allocate and fill the tuple lists, and it requires knowledge of how many entries will appear in the matrix, or at least a good upper bound, before the matrix is constructed. It is slightly faster, creating the same matrix in 60 seconds, 51 seconds of which is spent in GrB\_Matrix\_build.

```
GrB_Index *I, *J;
double *X ;
int64_t s = ((make_symmetric) ? 2 : 1) * nedges + 1 ;
I = malloc (s * sizeof (GrB_Index)) ;
J = malloc (s * sizeof (GrB_Index));
X = malloc (s * sizeof (double
                                 ));
if (I == NULL || J == NULL || X == NULL)
{
    // out of memory
    if (I != NULL) free (I) :
    if (J != NULL) free (J) :
    if (X != NULL) free (X) :
   return (GrB_OUT_OF_MEMORY) ;
int64_t ntuples = 0 ;
for (int64_t k = 0 ; k < nedges ; k++)
    GrB_Index i = simple_rand_i ( ) % nrows ;
    GrB_Index j = simple_rand_i ( ) % ncols ;
    if (no_self_edges && (i == j)) continue ;
    double x = simple_rand_x ( ) ;
    // A (i,j) = x
    I [ntuples] = i ;
    J [ntuples] = j ;
    X [ntuples] = x ;
   ntuples++;
    if (make_symmetric)
        // A (j,i) = x
        I [ntuples] = j ;
        J [ntuples] = i ;
        X [ntuples] = x ;
        ntuples++;
    }
GrB_Matrix_build (A, I, J, X, ntuples, GrB_SECOND_FP64) ;
```

The equivalent sprandsym function in MATLAB takes 150 seconds, but sprandsym uses a much higher-quality random number generator to create the tuples [I,J,X]. Considering just the time for sparse(I,J,X,n,n) in sprandsym (equivalent to GrB\_Matrix\_build), the time is 70 seconds. That is, each of these three methods, setElement and build in Suite-Sparse:GraphBLAS, and sparse in MATLAB, are equally fast.

## 6.4 Creating a finite-element matrix

Suppose a finite-element matrix is being constructed, with k=40,000 finite-element matrices, each of size 8-by-8. The following operations (in pseudo-MATLAB notation) are very efficient in SuiteSparse:GraphBLAS.

```
A = sparse (m,n) ; % create an empty n-by-n sparse GraphBLAS matrix
for i = 1:k
    construct a 8-by-8 sparse or dense finite-element F
    I and J define where the matrix F is to be added:
    I = a list of 8 row indices
    J = a list of 8 column indices
    % using GrB_assign, with the 'plus' accum operator:
    A (I,J) = A (I,J) + F
end
```

If this were done in MATLAB or in GraphBLAS with blocking mode enabled, the computations would be extremely slow. This example is taken from Loren Shure's blog on MATLAB Central, Loren on the Art of MATLAB [Dav07], which discusses the built-in wathen function. In MATLAB, a far better approach is to construct a list of tuples [I,J,X] and to use sparse(I,J,X,n,n). This is identical to creating the same list of tuples in GraphBLAS and using the GrB\_Matrix\_build, which is equally fast. The difference in time between using sparse or GrB\_Matrix\_build, and using submatrix assignment with blocking mode (or in MATLAB which does not have a nonblocking mode) can be extreme. For the example matrix discussed in [Dav07], using sparse instead of submatrix assignment in MATLAB cut the run time of wathen from 305 seconds down to 1.6 seconds.

In SuiteSparse:GraphBLAS, the performance of both methods is essentially identical, and roughly as fast as sparse in MATLAB. Inside SuiteSparse:GraphBLAS, GrB\_assign is doing the same thing. When performing A(I,J)=A(I,J)+F, if it finds that it cannot quickly insert an update into the A matrix, it creates a list of pending tuples to be assembled later on. When the matrix is ready for use in a subsequent GraphBLAS operation (one that normally cannot use a matrix with pending computations), the tuples are assembled all at once via GrB\_Matrix\_build.

GraphBLAS operations on other matrices have no effect. Thus, any GraphBLAS method or operation can be used to construct the F matrix in the example above, without affecting when the pending updates to A are completed.

The MATLAB wathen.m script is part of Higham's gallery of matrices [Hig02]. It creates a finite-element matrix with random coefficients for a 2D mesh of size nx-by-ny, a matrix formulation by Wathen [Wat87]. The pattern of the matrix is fixed; just the values are randomized. The GraphBLAS equivalent can use either GrB\_Matrix\_build, or GrB\_assign. Both methods have good performance. The GrB\_Matrix\_build version below is about 15% to 20% faster than the MATLAB wathen.m function, regardless of the problem size. It uses the identical algorithm as wathen.m.

```
int64_t ntriplets = nx*ny*64 ;
I = malloc (ntriplets * sizeof (int64_t));
J = malloc (ntriplets * sizeof (int64_t));
X = malloc (ntriplets * sizeof (double )) ;
if (I == NULL || J == NULL || X == NULL)
    FREE_ALL ;
    return (GrB_OUT_OF_MEMORY) ;
}
ntriplets = 0 ;
for (int j = 1; j \le ny; j++)
   for (int i = 1; i <= nx; i++)
        nn [0] = 3*j*nx + 2*i + 2*j + 1;
        nn [1] = nn [0] - 1;
       nn [2] = nn [1] - 1;
        nn [3] = (3*j-1)*nx + 2*j + i - 1;
        nn [4] = 3*(j-1)*nx + 2*i + 2*j - 3;
        nn [5] = nn [4] + 1;
        nn [6] = nn [5] + 1;
        nn [7] = nn [3] + 1;
        for (int krow = 0; krow < 8; krow++) nn [krow]--;
        for (int krow = 0; krow < 8; krow++)
        {
            for (int kcol = 0; kcol < 8; kcol++)
            {
               I [ntriplets] = nn [krow] ;
                J [ntriplets] = nn [kcol] ;
               X [ntriplets] = em (krow,kcol) ;
               ntriplets++ ;
            }
       }
   }
}
```

```
// A = sparse (I,J,X,n,n) ;
GrB_Matrix_build (A, I, J, X, ntriplets, GrB_PLUS_FP64) ;
```

The GrB\_assign version has the advantage of not requiring the user application to construct the tuple list, and is almost as fast as using GrB\_Matrix\_build. The code is more elegant than either the MATLAB wathen.m function or its GraphBLAS equivalent above. Its performance is comparable with the other two methods, but slightly slower, being about 5% slower than the MATLAB wathen, and 20% slower than the GraphBLAS method above.

```
GrB_Matrix_new (&F, GrB_FP64, 8, 8);
for (int j = 1; j \le ny; j++)
    for (int i = 1 ; i <= nx ; i++)
        nn [0] = 3*j*nx + 2*i + 2*j + 1;
        nn [1] = nn [0] - 1;
        nn [2] = nn [1] - 1;
        nn [3] = (3*j-1)*nx + 2*j + i - 1;
        nn [4] = 3*(j-1)*nx + 2*i + 2*j - 3;
        nn [5] = nn [4] + 1;
       nn [6] = nn [5] + 1;
       nn [7] = nn [3] + 1;
        for (int krow = 0; krow < 8; krow++) nn [krow]--;</pre>
        for (int krow = 0; krow < 8; krow++)
        {
            for (int kcol = 0; kcol < 8; kcol++)
                // F (krow,kcol) = em (krow, kcol)
               GrB_Matrix_setElement (F, em (krow,kcol), krow, kcol);
            }
        // A (nn,nn) += F
        GrB_assign (A, NULL, GrB_PLUS_FP64, F, nn, 8, nn, 8, NULL) ;
   }
}
```

Since there is no Mask, and since GrB\_REPLACE is not used, the call to GrB\_assign in the example above is identical to GxB\_subassign. Either one can be used, and their performance would be identical.

Refer to the wathen.c function in the Demo folder, which uses GraphBLAS to implement the two methods above, and two additional ones.

## 6.5 Reading a matrix from a file

The read\_matrix function in the Demo reads in a triplet matrix from a file, one line per entry, and then uses GrB\_Matrix\_build to create the matrix. It creates a second copy with GrB\_Matrix\_setElement, just to test that method and compare the run times. A comparison of build versus setElement has already been discussed in Section 6.3.

The function can return the matrix as-is, which may be rectangular or unsymmetric. If an input parameter is set to make the matrix symmetric,  $read_matrix$  computes  $A=(A+A^2)/2$  if A is square (turning all directed edges into undirected ones. If A is rectangular, it creates a bipartite graph, which is the same as the augmented matrix,  $A = [0 \ A \ ; \ A^2 \ 0]$  in pseudo-MATLAB notation.

If C is an n-by-n matrix, then C=(C+C')/2 can be computed as follows in GraphBLAS, (the scale2 function divides an entry by 2):

```
GrB_Descriptor_new (&dt2) ;
GrB_Descriptor_set (dt2, GrB_INP1, GrB_TRAN) ;
GrB_Matrix_new (&A, GrB_FP64, n, n) ;
GrB_eWiseAdd (A, NULL, NULL, GrB_PLUS_FP64, C, C, dt2) ; // A=C+C'
GrB_free (&C) ;
GrB_Matrix_new (&C, GrB_FP64, n, n) ;
GrB_UnaryOp_new (&scale2_op, scale2, GrB_FP64, GrB_FP64) ;
GrB_apply (C, NULL, NULL, scale2_op, A, NULL) ; // C=A/2
GrB_free (&A) ;
GrB_free (&scale2_op) ;
```

This is of course not nearly as elegant as A=(A+A')/2 in MATLAB, but with minor changes it can work on any type and use any built-in operators instead of PLUS, or it can use any user-defined operators and types. The above code in SuiteSparse:GraphBLAS takes 0.60 seconds for the Freescale2 matrix, slightly slower than MATLAB (0.55 seconds).

Constructing the augmented system is more complicated because Graph-BLAS does not yet have a simple way of specifying a range of row and column indices, as in A(10:20,30:50) in MATLAB. The application must instead build a list of indices first, I=[10, 11...20]. GraphBLAS does have a way of specifying all indices via I=GrB\_ALL, which results in A(:), but no easy way to specify a contiguous subset of indices. Thus, the following index lists I and J must first be constructed:

```
int64_t n = nrows + ncols ;
I = malloc (nrows * sizeof (int64_t)) ;
J = malloc (ncols * sizeof (int64_t)) ;
// I = 0:nrows-1
// J = nrows:n-1
if (I == NULL || J == NULL)
{
    if (I != NULL) free (I) ;
    if (J != NULL) free (J) ;
    return (GrB_OUT_OF_MEMORY) ;
}
for (int64_t k = 0 ; k < nrows ; k++) I [k] = k ;
for (int64_t k = 0 ; k < ncols ; k++) J [k] = k + nrows ;</pre>
```

Once the index lists are generated, however, the resulting GraphBLAS operations are fairly straightforward, computing A=[0 C; C'0].

```
GrB_Descriptor_new (&dt1) ;
GrB_Descriptor_set (dt1, GrB_INPO, GrB_TRAN) ;
GrB_Matrix_new (&A, GrB_FP64, n, n) ;
// A (nrows:n-1, 0:nrows-1) = C'
GrB_assign (A, NULL, NULL, C, J, ncols, I, nrows, dt1) ;
// A (0:nrows-1, nrows:n-1) = C
GrB_assign (A, NULL, NULL, C, I, nrows, J, ncols, NULL) ;
```

This takes 1.38 seconds for the Freescale2 matrix, almost as fast as A=[sparse(m,m) C; C' sparse(n,n)] in MATLAB (1.25 seconds).

Both calls to GrB\_assign use no accumulator, so the second one causes the partial matrix A=[0 0; C'0] to be built first, followed by the final build of A=[0 C; C'0]. A better method, but not an obvious one, is to use the GrB\_FIRST\_FP64 accumulator for both assignments. An accumulator enables SuiteSparse:GraphBLAS to determine that that entries created by the first assignment cannot be deleted by the second, and thus it need not force completion of the pending updates prior to the second assignment.

Any operator will suffice because it is not actually applied. An operator is only applied to the set intersection, and the two assignments do not overlap. If an accum operator is used, only the final matrix is built, and the time in GraphBLAS drops slightly to 1.25 seconds. This is a very small improvement because in this particular case, SuiteSparse:GraphBLAS is able to detect that no sorting is required for the first build, and the second one is a simple concatenation. In general, however, allowing GraphBLAS to postpone pending updates can lead to significant reductions in run time.

## 6.6 Triangle counting

A triangle in an undirected graph is a clique of size three: three nodes i, j, and k that are all pairwise connected. There are many ways of counting the number of triangles in a graph. Let A be a symmetric matrix with values 0 and 1, and no diagonal entries; this matrix is the adjacency matrix of the graph. Let E be the edge incidence matrix with exactly two 1's per column. A column of E with entries in rows i and j represents the edge (i, j) in the graph, A(i,j)=1 where i < j. Let E and E be the strictly lower and upper triangular parts of E, respectively.

The methods are listed in the table below. Most of them use a form of masked matrix-matrix multiplication. The methods are implemented in MATLAB in the tricount.m file, and in GraphBLAS in the tricount.c file, both in the GraphBLAS/Demo folder. Refer to the comments in those two files for details and derivations on how these methods work.

When a mask is present and not complemented, GrB\_INPO is GrB\_TRAN, and GrB\_INP1 is GxB\_DEFAULT, the SuiteSparse:GraphBLAS implementation of GrB\_mxm always uses a dot-product formulation. Thus, the  $\mathbf{C}\langle\mathbf{L}\rangle=\mathbf{U'L}$  method uses dot products. This provides a mechanism for the enduser to select a masked dot product matrix multiplication method in SuiteSparse:GraphBLAS, which is occassionally faster than the outer product method.

Each method is followed by a reduction to a scalar, via GrB\_reduce in GraphBLAS or by nnz or sum(sum(...)) in MATLAB.

method and	in MATLAB	in GraphBLAS
citation		
minitri [WBS15]	nnz(A*E==2)/3	$\mathbf{C} = \mathbf{AE},  ext{ then GrB\_apply}$
Burkhardt [Bur16]	$sum(sum((A^2).*A))/6$	$\mathbf{C}\langle\mathbf{A} angle=\mathbf{A}^2$
Cohen [ABG15, Coh09]	sum(sum((L*U).*A))/2	$\mathbf{C}\langle\mathbf{A} angle=\mathbf{L}\mathbf{U}$
Sandia [WDB <sup>+</sup> 17]	sum(sum((U*U).*U))	$\mathbf{C}\langle\mathbf{U}\rangle = \mathbf{U}\mathbf{U}$ (outer product)
SandiaDot	sum(sum((U'*L).*L))	$\mathbf{C}\langle\mathbf{L}\rangle = \mathbf{U}'\mathbf{L} \text{ (dot product)}$
SandiaL	sum(sum((L*L).*L))	$\mathbf{C}\langle\mathbf{L}\rangle = \mathbf{L}\mathbf{L}$ (outer product)

In general, the Sandia methods are the fastest of the 6 methods when implemented in GraphBLAS. The method in the KokkosKernels paper uses (L\*L).\*L via a masked matrix multiplication, but KokkosKernels stores its matrices in compressed sparse row form. GraphBLAS and MATLAB both store their matrices in compressed sparse column form, so the Sandia method is identical to (U\*U).\*U in MATLAB and  $\mathbf{C}\langle\mathbf{U}\rangle=\mathbf{U}\mathbf{U}$  in GraphBLAS. The

SandiaDot and SandiaL methods do not appear in [WDB<sup>+</sup>17], but they are named this way because they are simple extensions of the Sandia method.

The methods in MATLAB are slow because the matrix product is formed and then its entries are pruned via the element-wise multiplication (.\*). By contrast, <code>GrB\_mxm</code> only computes the entries residing in the mask, saving time and memory. This optimization is only exploited if the mask present and not complemented. Since the <code>minitri</code> method does not use a mask, its implementation in <code>GraphBLAS</code> has the same performance and memory requirements as the MATLAB version <code>nnz(A\*E==2)/3</code>. That is, both are very slow.

Performance results are shown in the following two tables. The first table is a list of matrices from the SuiteSparse Matrix Collection [DH11], listing the matrix name, the number of rows and columns, the number of edges in the graph, and the number of triangles. The matrices were symmetrized first with A=A+A' and the diagonal entries were removed. The first table splits into two sets. The first set of matrices also appear in the results from the Kokkos triangles paper [WDB+17].

The next table gives performance results on these matrices, with four methods. For each method, the run time in seconds and the rate is given, where the rate is the number of edges in the graph divided by the run time (listed in millions of edges per second). The first three methods in the table are for MATLAB and the two GraphBLAS methods, on a MacBook Pro (Retina, 13inch, Late 2013), 2.8 Ghz Intel Core i7, 16 GB RAM, OSX 10.11.6, MATLAB 2017a, with the clang 8.0.0 compiler. Only a single core was used for these results. In addition, the matrix L=tril(A) and/or U=triu(A) are used as-is without any reordering. The run times include the time to construct L or U. MATLAB failed on one matrix because U\*U is too large. For the first set of matrices, the outer product formulation ( $\mathbf{C}\langle\mathbf{U}\rangle=\mathbf{U}\mathbf{U}$ ) is always faster than the dot product formulation, but this is not the case for the second set.

The last column (Kokkos) is copied directly from [WDB<sup>+</sup>17]. The Kokkos results are from their implementation of sum(sum((L\*L).\*L)) using a masked sparse matrix-matrix multiply in KokkosKernels. These results were done on an Intel Xeon Haswell (E5-2698v3, 2.3GHz), with 512 GB RAM, 32 cores and 2 hyperthreads per core, using the Intel icc 17.1 compiler. Unlike the other three methods, L is sorted by decreasing row degree, which improves the performance. The Kokkos time includes the time taken to do the sort. The run time listed is the best time obtained from several runs with 1 to 32

threads.

Comparing GraphBLAS and Kokkos is difficult since these results were obtained on different machines. Also, the results in [WDB+17] provide just the best-obtained parallel results, not the results on a single core. In addition, these results are with a reordered L in Kokkos, but not in GraphBLAS. Wolf et al. [WDB+17] state that reordering L improves the run time. However, with these many caveats, the last column lists the speedup of Kokkos over the GraphBLAS outer-product formulation. Since the Kokkos method is parallel these preliminary comparisons indicate that the sequential performance of GraphBLAS is competitive. Using up to 32 threads, Kokkos is about 3 to 18 faster than SuiteSparse:GraphBLAS, which is currently sequential (median speedup of about 9). Further comparisons are required, however. A parallel implementation of the matrix-matrix multiply in GrB\_mxm is also in progress.

matrix	n	# edges	# triangles
SNAP/cit-HepPh	34,546	420,877	1,276,868
SNAP/cit-HepTh	27,770	$352,\!285$	$1,\!478,\!735$
SNAP/email-EuAll	$265,\!214$	$364,\!481$	$267,\!313$
SNAP/soc-Epinions1	75,888	405,740	1,624,481
SNAP/soc-Slashdot0811	77,360	$469,\!180$	551,724
SNAP/soc-Slashdot0902	82,168	$504,\!230$	$602,\!592$
SNAP/amazon0312	400,727	2,349,869	3,686,467
SNAP/amazon0505	$410,\!236$	2,439,437	3,951,063
SNAP/amazon0601	403,394	2,443,408	3,986,507
SNAP/cit-Patents	3,774,768	16,518,947	7,515,023
SNAP/soc-LiveJournal1	$4,\!847,\!571$	42,851,237	285,730,264
Gleich/wb-edu	$9,\!845,\!725$	$46,\!236,\!105$	254,718,147
SNAP/p2p-Gnutella09	8,115	26,013	2,354
Mallya/lhr71	70,304	1,492,794	$160,\!592$
Freescale/Freescale2	2,999,349	5,744,934	21,027,280
Freescale/circuit5M	5,558,326	26,983,926	31,019,473
DIMACS10/hugebubbles-00020	21,198,119	31,790,179	0
vanHeukelum/cage15	5,154,859	47,022,346	36,106,416

matrix	MATLAB		$\mathbf{C}\langle\mathbf{U} angle=\mathbf{L}'\mathbf{U}$		$\mathbf{C}\langle\mathbf{U} angle=\mathbf{U}\mathbf{U}$		Kokkos		
	time	rate	time	$_{\mathrm{rate}}$	time	rate	time	$_{\mathrm{rate}}$	speedup
SNAP/cit-HepPh	0.363	1.16	0.180	2.47	0.049	9.59	0.0044	79.9	8.3
SNAP/cit-HepTh	0.415	0.85	0.171	2.05	0.046	8.31	0.0050	72.5	8.7
SNAP/email-EuAll	1.264	0.29	0.133	2.73	0.035	10.33	0.0058	70.7	6.8
SNAP/soc-Epinions1	0.778	0.52	0.376	1.08	0.067	6.01	0.0039	108.0	18.0
SNAP/soc-Slashdot0811	0.990	0.47	0.318	1.47	0.052	9.04	0.0061	76.8	8.5
SNAP/soc-Slashdot0902	0.985	0.51	0.339	1.49	0.059	8.61	0.0063	80.1	9.3
SNAP/amazon0312	1.285	1.83	0.514	5.32	0.306	8.61	0.0754	30.7	3.6
SNAP/amazon0505	1.018	2.07	0.545	4.48	0.297	8.21	0.0177	133.0	16.2
SNAP/amazon0601	1.018	2.40	0.563	4.34	0.296	8.27	0.0184	132.0	16.0
SNAP/cit-Patents	11.026	1.50	4.416	3.74	2.300	7.18	0.4970	31.5	4.4
SNAP/soc-LiveJournal1	11.026	0.40	39.767	1.08	10.123	4.23	0.7330	58.5	13.8
Gleich/wb-edu	67.636	0.68	8.016	5.77	3.605	12.82	0.2320	199.0	15.5
SNAP/p2p-Gnutella09	0.004	6.50	0.002	10.65	0.001	24.24			
Mallya/lhr71	0.252	5.93	0.058	25.90	0.030	50.37			
Freescale/Freescale2	0.741	7.75	0.501	11.46	0.276	20.83			
Freescale/circuit5M	mem		2.819	9.57	194.142	0.14			
DIMACS10/hugebubbles-00020	7.406	4.29	3.417	9.30	6.568	4.84			
vanHeukelum/cage15	10.187	4.62	4.407	10.67	2.443	19.25			

The outer product  $\mathbf{C}\langle \mathbf{U} \rangle = \mathbf{U}\mathbf{U}$  in GraphBLAS is very simple:

```
int64_t ntriangles ;
GrB_Index n, one = 1 ;
GrB_Matrix C, U ;
GrB_Matrix_nrows (&n, A) ;

// U = triu (A, 1)
GrB_Matrix_new (&U, GrB_UINT32, n, n) ;
GxB_select (U, NULL, NULL, GxB_TRIU, A, &one, NULL) ;

// C<U> = U*U
GrB_Matrix_new (&C, GrB_UINT32, n, n) ;
GrB_mxm (C, U, NULL, GxB_PLUS_TIMES_UINT32, U, U, NULL) ;

// ntriangles = sum (C)
GrB_reduce (&ntriangles, NULL, GxB_PLUS_INT64_MONOID, C, NULL) ;

GrB_free (&C) ;
GrB_free (&U) ;
```

The dot product method  $\mathbf{C}\langle\mathbf{U}\rangle = \mathbf{L}'\mathbf{U}$  in GraphBLAS is similar:

```
int64_t ntriangles ;
GrB_Index n, one = 1, minusone = -1;
GrB_Matrix C, L, U;
GrB_Matrix_nrows (&n, A) ;
// U = triu (A, 1)
GrB_Matrix_new (&U, GrB_UINT32, n, n);
GxB_select (U, NULL, NULL, GxB_TRIU, A, &one, NULL) ;
// L = tril (A,-1)
GrB_Matrix_new (&L, GrB_UINT32, n, n);
GxB_select (L, NULL, NULL, GxB_TRIL, A, &minusone, NULL) ;
// C<U> = L'*U
GrB_Matrix_new (&C, GrB_UINT32, n, n);
GrB_Descriptor_new (&d) ;
GrB_Descriptor_set (d, GrB_INPO, GrB_TRAN) ;
GrB_mxm (C, U, NULL, GxB_PLUS_TIMES_UINT32, L, U, d) ;
GrB_free (&d) ;
// ntriangles = sum (C)
GrB_reduce (&ntriangles, NULL, GxB_PLUS_INT64_MONOID, C, NULL) ;
GrB_free (&C) ;
GrB_free (&L) ;
GrB_free (&U) ;
```

# 6.7 User-defined types and operators: double complex and struct-based

The Demo folder contains two working examples of user-defined types, first discussed in Section 4.1.1: double complex, and a user-defined typedef called wildtype with a struct containing a string and a 4-by-4 float matrix.

**Double Complex:** GraphBLAS does not have a native complex type, but this can be easily added as a user-defined type. The Complex\_init function in the usercomplex.c file in the Demo folder creates the Complex type based on the ANSI C11 double complex type.

GrB\_Type\_new (&Complex, double complex) ;

Next, it creates a full suite of operators that correspond to every built-in GraphBLAS operator, both binary and unary. In addition, it creates the operators listed in the following table, where D is double and C is Complex.

name	types	MATLAB	description		
		equivalent			
Complex_complex	$D \times D \to C$	z=complex(x,y)	complex from real and imag.		
Complex_conj	$C \to C$	z=conj(x)	complex conjugate		
Complex_real	$C \to D$	z=real(x)	real part		
Complex_imag	$C \to D$	z=imag(x)	imaginary part		
Complex_angle	$C \to D$	z=angle(x)	phase angle		
Complex_complex_real	$D \to C$	z=complex(x,0)	real to complex real		
Complex_complex_imag	$D \to C$	z=complex(0,x)	real to complex imag.		

The Complex\_init function creates two monoids (Complex\_add\_monoid and Complex\_times\_monoid) and a semiring Complex\_plus\_times that corresponds to the conventional linear algebra for complex matrices. The include file usercomplex.h in the Demo folder is available so that this user-defined Complex type can easily be imported into any other user application. When the user application is done, the Complex\_finalize function frees the Complex type and its operators, monoids, and semiring.

Struct-based: In addition, the wildtype.c program creates a user-defined typedef of a struct containing a dense 4-by-4 float matrix, and a 64-character string. It constructs an additive monoid that adds two 4-by-4 dense matrices, and a multiplier operator that multiplies two 4-by-4 matrices. Each of these 4-by-4 matrices is treated by GraphBLAS as a "scalar" value, and they can be manipulated in the same way any other GraphBLAS type can be manipulated. The purpose of this type is illustrate the endless possibilities of user-defined types and their use in GraphBLAS.

# 7 Installing SuiteSparse:GraphBLAS

GraphBLAS makes extensive use of features in the ANSI C11 standard, and thus a C compiler supporting this version of the C standard is required. On the Mac (OS X), clang 8.0.0 in Xcode version 8.2.1 is sufficient, although earlier versions of Xcode may work as well. For the GNU gcc compiler, version 4.9 or later is required. For the Intel icc compiler, version 18.0 or later is required. Version 2.8.12 or later of cmake is required; version 3.0.0 is preferred.

To compile SuiteSparse:GraphBLAS and the demo programs, simply type make in the main GraphBLAS folder, which compiles the library and runs several demos.

GraphBLAS is not yet parallel, but it is thread-safe if multiple simultaneous calls are made to GraphBLAS functions. For this usage, GraphBLAS must be compiled with OpenMP so that GraphBLAS has access to a critical section mechanism. OpenMP is optional if the user application does not make multiple simultaneous calls to GraphBLAS.

If cmake or make fail, it might be that your default compiler does not support ANSI C11. Try another compiler. For example, try one of these options. Go into the build directory and type:

```
CC=gcc cmake ..

CC=gcc-6 cmake ..

CC=xlc cmake ..

CC=icc cmake ..
```

Then do make in the build directory. If this still fails, see the CMakeLists.txt file. You may need to pass compiler-specific options to your compiler. Locate this section in the CMakeLists.txt file. Use the set command in cmake, as in the example below, to set the compiler flags you need.

```
# check which compiler is being used. If you need to make
# compiler-specific modifications, here is the place to do it.
if ("${CMAKE_C_COMPILER_ID}" STREQUAL "GNU")
    # cmake 2.8 workaround: gcc needs to be told to do ANSI C11.
    # cmake 3.0 doesn't have this problem.
    set (CMAKE_C_FLAGS "-std=c11 -lm -fopenmp")
    ...
elseif ("${CMAKE_C_COMPILER_ID}" STREQUAL "Intel")
```

```
elseif ("${CMAKE_C_COMPILER_ID}" STREQUAL "Clang")
...
elseif ("${CMAKE_C_COMPILER_ID}" STREQUAL "MSVC")
...
endif ( )
```

Once cmake and make finish, run the demos in the GraphBLAS/Demo folder:

```
cd ../Demo
./demo
```

The ./demo command is a script that runs the demos with various input matrices in the Demo/Matrix folder. The output of the demos will be compared with expected output files in Demo/Output.

To install the library in /usr/local/lib and /usr/local/include, go to the top-level GraphBLAS folder and type:

```
sudo make install
```

Several compile-time options can be selected by editing the Source/GB.h file, but these are meant only for code development of SuiteSparse:GraphBLAS itself, not for end-users of SuiteSparse:GraphBLAS.

To perform the extensive tests in the Test folder, and the statement coverage tests in Tcov, MATLAB R2017A is required. See the README.txt files in those two folders for instructions on how to run the tests.

To remove all compiled files, type make distclean in the top-level Graph-BLAS folder.

NOTE: SuiteSparse:GraphBLAS has not yet been ported to Windows. However, with cmake the port to Windows should be straightforward (this is in progress).

# 8 Acknowledgements

I would like to thank Jeremy Kepner (MIT Lincoln Laboratory Supercomputing Center), and the GraphBLAS API Committee: Aydın Buluç (Lawrence Berkeley National Laboratory), Timothy G. Mattson (Intel Corporation) Scott McMillan (Software Engineering Institute at Carnegie Mellon University), José Moreira (IBM Corporation), and Carl Yang (UC Davis), for creating the GraphBLAS specification and for patiently answering my many questions while I was implementing it.

I would also like to thank Sébastien Villemot (Debian Developer, http://sebastien.villemot.name) for helping me with various build issues and other code issues with GraphBLAS (and all of SuiteSparse) for its packaging in Debian Linux.

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Provides a basic overview of many sparse matrix algorithms and a simple sparse matrix data structure. The sparse data structure used in the book is much like the one in both MATLAB and SuiteSparse:GraphBLAS. A series of 42 lectures are available on YouTube; see the link at http://faculty.cse.tamu.edu/davis/publications.html. DOI: https://dx.doi.org/10.1137/1.9780898718881

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Abstract: Wilkinson defined a sparse matrix as one with enough zeros that it pays to take advantage of them. This informal yet practical definition captures the essence of the goal of direct methods for solving sparse matrix problems. They exploit the sparsity of a matrix to solve problems economically: much faster and using far less memory than if all the entries of a matrix were stored and took part in explicit computations. These methods form the backbone of a wide range of problems in computational science. A glimpse of the breadth of applications relying on sparse solvers can be seen in the origins of matrices in published matrix benchmark collections (Duff and Reid 1979a, Duff, Grimes and Lewis 1989a, Davis and Hu 2011). The goal of this survey article is to impart a working knowledge of the underlying theory and practice of sparse direct methods for solving linear systems and least-squares problems, and to provide an overview of the algorithms, data structures, and software available to solve these problems, so that the reader can both understand the methods and know how best to use them. DOI: https://dx.doi.org/10.1017/S0962492916000076

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From the preface: Graphs are among the most important abstract data types in computer science, and the algorithms that operate on them are critical to modern life. Graphs have been shown to be powerful tools for modeling complex problems because of their simplicity and generality. Graph algorithms are one of the pillars of mathematics, informing research in such diverse areas as combinatorial optimization, complexity theory, and topology. Algorithms on graphs are applied in many ways in today's worldfrom Web rankings to metabolic networks, from finite element

meshes to semantic graphs. The current exponential growth in graph data has forced a shift to parallel computing for executing graph algorithms. Implementing parallel graph algorithms and achieving good parallel performance have proven difficult. This book addresses these challenges by exploiting the well-known duality between a canonical representation of graphs as abstract collections of vertices and edges and a sparse adjacency matrix representation. This linear algebraic approach is widely accessible to scientists and engineers who may not be formally trained in computer science. The authors show how to leverage existing parallel matrix computation techniques and the large amount of software infrastructure that exists for these computations to implement efficient and scalable parallel graph algorithms. The benefits of this approach are reduced algorithmic complexity, ease of implementation, and improved performance. DOI: https://doi.org/10.1137/1.9780898719918

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Triangle counting serves as a key building block for a set of important graph algorithms in network science. In this paper, we address the IEEE HPEC Static Graph Challenge problem of triangle counting, focusing on obtaining the best parallel performance on a single multicore node. Our implementation uses a linear algebra-based approach to triangle counting that has grown out of work related to our miniTri

data analytics miniapplication and our efforts to pose graph algorithms in the language of linear algebra. We leverage KokkosKernels to implement this approach efficiently on multicore architectures. Our performance results are competitive with the fastest known graph traversal-based approaches and are significantly faster than the Graph Challenge reference implementations, up to 670,000 times faster than the C++ reference and 10,000 times faster than the Python reference on a single Intel Haswell node.