

# Documentation for Zn.h and Zn.c

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## Header

```
#ifndef __Zn_h
#define __Zn_h

#include <stdlib.h>
#include <stdio.h>
#define allocZV(n) ((int *) calloc(n,sizeof(int)))
#define freeZV(a) free(a)

int *setstdZV(int *c,int n,int k);
int *printZV(int *c,int n);
char *Zn_vect2csvstring(int *c,int n,char *string);
int *fprintZV(FILE *stream,int *c,int n);
int minZV(int *a,int n);
int maxZV(int *a,int n);
int *copyZV(int *a,int *c,int n);
int *sumZV(float ax,int *a,float bx,int *b,int *c,int n);
int *deriv1ZV(int *a,int *c,int n);
int *deriv2ZV(int *a,int *c,int n);
int productZV(int *a,int n);
int intfindZV(int *a,int n,int i);
int indx2addZV(int *indx,int *dim,int rank);
int *add2indxZV(int add,int *indx,int *dim,int rank);
int nextaddZV(int add,int *indx1,int *indx2,int *dim,int rank);
int indx2add3ZV(int *indx,int rank);
int *add2indx3ZV(int add,int *indx,int rank);
int neighborZV(int *indx,int *c,int *dim,int rank,int type,int *wrap,int *mid);
int Zn_sameset(int *a,int *b,int *work,int n);
void Zn_sort(int *a,int *b,int n);
int Zn_issort(int *a,int n);

/***** combinatorics *****/

int Zn_permute(int *a,int *b,int n,int k);
int Zn_permutelex(int *seq,int n);

#endif
```

Requires: <stdio.h>, <stdlib.h>, "random.h"

Example program: none

Written 10/01; some testing. Written on Linux, copied to Macintosh. Added tensor routines 10/31/01. Added fprintZV and intfindZV 3/03. Modified neighborZV and added add2indx3ZV and indx2add3ZV 7/7/03. Added nextaddZV 7/14/03. Added Zn\_vect2csvstring 1/11/08. Added Zn\_sameset 2/15/08. Added Zn\_permutelex 1/30/12.

This library is comparable to the Rn.c and Cn.c libraries, but is designed for vectors of integers rather than real or complex numbers. There are also some routines for keeping track of the indicies of tensors, routines which can also be used for converting the base of a number. For all routines where they are used, a, b, and c are pointers to arrays of integers where c is the output array, and n is the number of elements (numbered from 0 to n-1). Unless otherwise specified, the functions return c to allow easy concatenation of routines. Routines don't have internal error checking and assume the input is good (all vectors defined and  $n \geq 1$ ).

The tensor routines assume a tensor is stored sequentially in memory with the 0'th dimension varying most slowly, and the *rank*-1'th dimension varying most quickly. For example, a matrix with *m* rows and *n* columns (see Rn.c), which is stored with values filling one row before starting the next row, would be interpreted as having *m* as the 0'th dimension and *n* as the first dimension. In that case, the address of index *i,j* is  $a = n * i + j$  and the index of address *a* is  $i = a / n, j = a \% n$ , where / represents integer arithmetic and % is the modulus operator. These routines can also be used to convert a number from one base to another. In this case all the dimension values are set to the base number, the rank is the number of digits, and the 0'th digit is the most significant digit.

			dim[1]=4			
			<u>0</u>	<u>1</u>	<u>2</u>	<u>3</u>
dim[0]=3	0		0	1	2	3
	1		4	5	6	7
	2		8	9	10	11

```
wptype[j]=wrap[i]>>2*j&3;
```

The result is 0, 1, or 2 on each dimension, where 0 indicates no wrap-around, 1 indicates wrap-around towards the low side, and 2 indicates wrap-around towards the high side.

## Functions

### setstdZV

Initializes a vector, where the initial value depends on k. k=0 yields all 0's, k=1 yields all 1's, k<0 yields all zeros except for the element at position -k which is 1, k=2 yields sequential numbering from 0 to n-1, and k=3 yields either a 0 or 1 for each position, chosen randomly with equal probability. It's not defined for k>3.

### printZV

Prints the vector contents in a single row of text.

```
char *Zn_vect2csvstring(int *c,int n,char *string);
```

This prints the vector contents of c, which has n elements, to the string string as a comma-separated vector without a trailing newline. No check is made that the string is long enough for the result. The string is returned.

### fprintZV

Identical to printZV, but prints to filestream stream.

### minZV

Returns the smallest value in the vector.

### maxZV

Returns the largest value in the vector.

### copyZV

Copies a vector.

### deriv1ZV

Returns the first derivative of a vector, assuming unit spacing. n must be at least 3. See documentation for Rn.c for more details. The function requires a division by two for the calculation of each element of c; as these are vectors of integers and integer arithmetic truncates any fractional part, this means that the derivative will underestimate the actual derivative by about 0.5, on average. The derivative of 0,1,2,3 is 1,1,1,1 as it should be, but the derivative of 0,0,1,1 is 0,0,0,0. The second derivative function has no division and so does not have round-off problems.

### deriv2ZV

Returns the second derivative of a vector, assuming unit spacing. n must be at least 3. See documentation for Rn.c for more details. The function has no division and so does not have round-off problems.

### productZV

Calculates the product of all the elements of a. It's mostly useful for determining the maximum address of a tensor or the largest number expressible in a certain base, both of which are described below.

### intfindZV

Looks for the value  $i$  in the vector  $a$ , which has size  $n$ , returning the index where  $i$  is found if it is found, and  $-1$  if it is not found.

#### indx2addZV

Converts the index of a tensor element to its address, assuming values are numbered sequentially.  $indx$  and  $dim$  have rank elements each, where  $indx$  is the index and  $dim$  is the size of each tensor dimension. Negative values are permitted in  $indx$ , which can be useful for determining relative addresses, although all values of  $dim$  should be  $\geq 1$ . This does not check if the address is less than 0 or greater than the tensor size. See discussion below.

#### add2indxZV

Converts an address to the index. The address must be positive. This does not check if the input address is possible. It returns  $indx$ . See discussion below.

#### nextaddZV

Inputs an address as  $add$  and two indicies as  $indx1$  and  $indx2$ , as well as the tensor dimensionality and rank in  $dim$  and  $rank$ , respectively. The return value is the next sequential address which is within the sub-tensor defined with the lower corner at  $indx1$  and upper corner at  $indx2$ . This is particularly useful in a for loop, in which it is desirable to look at all addresses in a certain region of the tensor. For example,

```
add1=indx2addZV(indx1,dim,rank);
add2=indx2addZV(indx2,dim,rank);
for(add=add1;add<=add2;add=nextaddZV(add,indx1,indx2,dim,rank));
```

This function assumes that every element of  $indx1$  is less than or equal to the respective element of  $indx2$ , and it does not account for periodic boundaries. The return value is undefined if  $add$  is not within the range defined by  $indx1$  and  $indx2$ . If  $add$  corresponds to  $indx2$ , the returned value is  $add+1$ .

#### indx2add3ZV

Identical to  $indx2addZV$ , except that all  $dim$  elements are set equal 3.

#### add2indx3ZV

Identical to  $add2indxZV$ , except that all  $dim$  elements are set equal 3.

#### neighborZV

Returns the number of neighbors of an element along with their addresses, sent back in  $c$ . A return value of  $-1$  means that temporary memory could not be allocated. Otherwise there is no error checking.  $indx$  is the element and  $dim$  and  $rank$  refer to the tensor.  $mid$  is used by the function to return the mid-point of  $c$ ; the neighbors from  $c[0]$  to  $c[mid-1]$  logically precede the indexed element, whereas  $c[mid]$  to  $c[n-1]$  logically follow the indexed element.  $mid$  may be sent in as NULL if this information is not wanted. There are several allowable type values:

<u>type</u>	<u>neighbor</u>	<u>wrap-around</u>	<u>order</u>	<u>size of c</u>	<u>size of wrap</u>
-1	This code is used to free any temporary memory that was allocated				
0	nearest	none	increasing	$\leq 2*rank$	N/A
1	nearest	all	none	$\leq 2*rank$	N/A
2	all	none	increasing	$\leq 3^{rank}-1$	N/A
3	all	all	none	$\leq 3^{rank}-1$	N/A
4	nearest	partial	none	$\leq 2*rank$	rank
5	all	partial	none	$\leq 3^{rank}-1$	rank

6	nearest	partial	none	$\leq 2^{\text{rank}}$	$2^{\text{rank}}$
7	all	partial	none	$\leq 3^{\text{rank}-1}$	$3^{\text{rank}-1}$

Nearest neighbors are just those above, below, left, right, etc. of an element; they share a side. All neighbors includes also diagonal and corner neighbors. Wrap-around refers to the neighbors of an edge element, which can be either nothing on the edge side, or the wrapped around element. The non-wrap-around routines return  $c$  in increasing order of addresses. For types 0 to 5, addresses are not repeated in  $c$ , even if a neighbor is a neighbor in multiple ways, although they are for types 6 and 7. The size of  $c$  needs to be allocated beforehand to be large enough for the result. Partial wrap-around means that some dimensions wrap-around, whereas others don't; this information is sent to the routine in the first  $\text{rank}$  elements of  $\text{wrap}$ , where a 0 means don't wrap and a 1 means wrap. For types 6 and 7, the information is overwritten with the wrap code, which tells in what way each neighbor is a neighbor. In the wrap code, pairs of bits are associated with each dimension (low order bits with low dimension), with the bits equal to 00 for no wrapping in that dimension, 01 for wrapping towards the low side, and 10 for wrapping towards the high side. See the example below.

Functions types 2, 3, 5, 6, and 7 require small amounts of temporary memory as scratch space. This is allocated the first time the function is called, but is not freed afterwards, allowing the function to be called multiple times with minimal run-time overhead. At the end, it is proper to call the function one more time, with a type value of  $-1$ , to free any memory that was allocated. In this case, all other parameters are ignored.

```
int Zn_sameset(int *a, int *b, int *work, int n);
```

Returns 1 if vector  $b$  is a permutation (or identical) to vector  $a$ , and 0 if the set of elements in  $b$  is different than that in vector  $a$ . Each vector has  $n$  elements.  $\text{work}$  is a required  $n$ -element vector that is simply used as work space.

```
void Zn_sort(int *a, int *b, int n);
```

Sorts integer vector  $a$  into vector  $b$ , each of which is of size  $n$ .  $a$  and  $b$  are allowed to be the same vector; or, set  $b$  to NULL and  $a$  will be sorted. This function is identical to the `RnSort.c` function `sortV`, but for integers.

```
int Zn_issort(int *a, int n);
```

Determines if  $a$ , which has size  $n$ , is sorted. Returns 1 if all members of  $a$  are equal to each other (which also applies if  $n$  is 0 or 1), 2 if  $a$  is increasing but not strictly increasing, 3 if  $a$  is strictly increasing, -2 if  $a$  is decreasing but not strictly decreasing, -3 if  $a$  is strictly decreasing, or 0 if  $a$  is unsorted such that it is neither increasing nor decreasing. (Strictly increasing means  $a_{i-1} < a_i$  for all  $i$ , while just increasing means  $a_{i-1} \leq a_i$  for all  $i$ ).

## Combinatorics

```
int Zn_permute(int *a, int *b, int n, int k);
```

Finds all distinct permutations of integer vectors that have up to 3 elements. Send in an integer vector in  $a$ , a destination vector in  $b$  (which is not allowed to be the same as  $a$ ), the size of the vectors in  $n$  (between 0 and 3, inclusive), and the permutation code in  $k$ . Entering 0 for  $k$  returns  $b$  equal to  $a$ , which is the first permutation in the sequence. The function return value is the next  $k$  value in the sequence, which will be between 0 and  $n!-1$ . Upon completion of the sequence, 0

is returned so that it will start over again. This accounts for duplicate values in  $a$  by always returning the code for the next distinct permutation – the return value is always 1 greater than  $k$ , modulo  $n$ , when each element is different from each other, but may skip ahead if there are non-distinct elements.

```
int Zn_permutelex(int *seq, int n);
```

This computes the next permutation of the items listed in `seq`, of which there are  $n$  items, according to lexicographical ordering, and puts the result back in `seq`. Multiple items of `seq` are allowed to equal each other; if this happens, then these items are not permuted (e.g. if the starting sequence is 1,2,2, then subsequent sequences are 2,1,2, and 2,2,1, which is the end). This returns 1 when the final sequence is reached, 2 when the sequence wraps around to the start, and 0 otherwise. If the final sequence is sent in as an input, then, this reverses the sequence so as to start over again. This algorithm is from the web and is supposedly from Dijkstra, 1997, p. 71.