Boundary operators on LAR *

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Abstract

The various versions of boundary operators on Linear Algebraic Representation of cellular complexes are developed in this module, in order to maintain under focus their proper development, including the possible special cases.

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1 Introduction

In the current LarLib implementation, we have to distinguish between between dimension-independent, dimension-dependent, oriented and non-oriented operators. Therefore a code refactoring of LarLib—related to boundary/coboundary operators—started here, with the aim of both providing a precise mathematical definition within the LAR framework, and to simplify and generalise the implemented algorithms.

2 Implementation

We start this section by making a distinction between the (matrices of) boundary operators for the linear spaces C_k of chains over the field $\mathbb{Z}_2 = \{0,1\}$ and over the field \mathbb{Z} of integer numbers. We call either non-oriented or oriented the corresponding boundary operators, respectively, since the matrix elements take values within the sets $\{-1,0,+1\}$ or $\{0,1\}$, correspondingly. Of course, the associated matrices of coboundary operators are their transpose matrices.

2.1 Non-oriented operators

For several computations, the knowledge of the matrices of non-oriented boundary operators is sufficient. Therefore we will use such tool wherever possible, since its computation is much faster in term of computing time.

In the following we provide be binary operator matrices provided by two implementations, respectively named boundary and larUnsignedBoundary2. The first one works correctly only with convex cells; the second one works also with non-convex but path-connected cells.

2.1.1 Dimension-independence

As we show in the following, in order to compute the non-oriented boundary operator ∂_d , it is sufficient to have knowledge of the M_d and M_{d-1} characteristic matrices of d-cells and their (d-1)-facets, at least in the case of cellular complexes with convex cells. Conversely, for more general non-convex but simply-connected cells, also the M_{d-2} matrix is needed.

Convex-cells The algorithm used is pretty easy to present. The compressed characteristic matrices of d-cells and (d-1)-cells, denoted as cells and facets, respectively, are first put in csr format as csrCV and csrFV. Then the incidence matrix csrFC in compressed sparse row format is computed by matrix product of the compressed characteristic matrices.

The element (i, j) of this matrix provides the number of vertices in the intersection of facet i and cell j, whereas the number of non-zero elements in each csrFV row gives the number of vertices of the facet represented by the row, and is stored in facetLengths.

The boundary function—to be used only with dimension-independent LAR convex cells—is written efficiently in the following script, by using only the standard functions and attributes of the scipy.sparse module.

The variable facetCoboundary stores in a list, for every facet (for h in range(m)) the list of cells in its *coboundary*, to be stored in the output csr_matrix boundary matrix as column indices of elements with non-zero (i.e. 1) value.

Notice that both the computation of facetCoboundary contents, and the output of the compressed boundary matrix, are performed in the most efficient way—according to the internal design of the scipy's csr sparse data structure.

```
\langle convex\text{-cells boundary operator } 3 \rangle \equiv
     """ convex-cells boundary operator --- best implementation """
     def larBoundary(cells,facets):
         lenV = max(max(CAT(cells)), max(CAT(facets)))+1
         csrCV = csrCreate(cells,lenV)
         csrFV = csrCreate(facets,lenV)
         csrFC = csrFV * csrCV.T
         facetLengths = [csrFacet.getnnz() for csrFacet in csrFV]
         m,n = csrFC.shape
         facetCoboundary = [[csrFC.indices[csrFC.indptr[h]+k]
              for k,v in enumerate(csrFC.data[csrFC.indptr[h]:csrFC.indptr[h+1]])
                  if v==facetLengths[h]] for h in range(m)]
         indptr = [0]+list(cumsum(AA(len)(facetCoboundary)))
         indices = CAT(facetCoboundary)
         data = [1]*len(indices)
         return csr_matrix((data,indices,indptr),shape=(m,n),dtype='b')
```

Fragment referenced in 29b.

2.2 Non-convex LAR cells

A more general larUnsignedBoundary2 operator is given in the following, aiming at compute the boundary matrix for general non-convex cellular decompositions, including multiply connected LAR models. Notice that in this case an input triple made by CV, FV, and EV is needed, where—more in general embedded in \mathbf{E}^d —they stand for the (binary compressed) characteristic matrices M_d , M_{d-1} , and M_{d-2} .

Boundary operator from 3-chains to 2-chains

Fragment referenced in 29b.

```
\langle path\text{-}connected\text{-}cells \ boundary \ operator \ 4 \rangle \equiv
     """ path-connected-cells boundary operator """
     def larUnsignedBoundary2(CV,FV,EV):
         out = larBoundary(CV,FV)
         def csrRowSum(h):
             return sum(out.data[out.indptr[h]:out.indptr[h+1]])
         unreliable = [h for h in range(len(FV)) if csrRowSum(h) > 2]
         if unreliable != []:
             csrBBMat = larBoundary(FV,EV) * larBoundary(CV,FV)
             lenV = max(max(CAT(CV)), max(CAT(FV)), max(CAT(EV)))+1
             FE = larcc.crossRelationO(lenV,FV,EV)
              out = csrBoundaryFilter2(unreliable,out,csrBBMat,CV,FE)
         return out
     def boundary3(CV,FV,EV):
         out = larUnsignedBoundary2(CV,FV,EV)
         lenV = max(max(CAT(CV)), max(CAT(FV)), max(CAT(EV)))+1
         VV = AA(LIST)(range(lenV))
         csrBBMat = scipy.sparse.csc_matrix(larBoundary(FV,EV) * larUnsignedBoundary2(CV,FV,EV))
         def csrColCheck(h):
              return any([val for val in csrBBMat.data[csrBBMat.indptr[h]:csrBBMat.indptr[h+1]] if v
         unreliable = [h for h in range(len(CV)) if csrColCheck(h)]
         if unreliable != []:
             FE = larcc.crossRelationO(lenV,FV,EV)
              out = csrBoundaryFilter3(unreliable,out,csrBBMat,CV,FE)
         return out
Fragment defined by 4, 5a.
```

Boundary operator from 2-chains to 1-chains First the boundary operator for the convex case is computed within the out variable of csr_matrix type. Then every out row (i.e. every (d-1)-facet of the d-complex) is tested for reliability, since every (d-1)-face can be shared by at most two d-cells in a d-complex. When this condition is not satisfied, deeper tests are needed to understand what row elements must be forced to value 1, since the (d-1)-face itself is a subset, but not actually a facet, of the corresponding d-cell.

In presence of some "unreliable" facets, the matrix csrBBMat of the operator $\partial_{d-1} \circ \partial_d$ and the relation FE between faces of dimensions d-1 and d-2 are computed. Now, let us notice that the columns of csrBBMat report the number of incidences of the d-2 faces (as belonging to (d-1)-facets embedded on the boundary) and d-cells (that are associated to

such matrix columns). Hence, in a regular (convex) d-complex, such numbers are always even, and in \mathbb{Z}_2 arithmetic are reduced to zero, in order to satisfy the fundaments equation $\partial \partial = 0$.

Conversely, with non-convex LAR cells, some incidence numbers may get odd values, due to the non-strict coincidence between cell facets and vertex subsets. Therefore, for "unreliable" h rows (facets) the csrBBMat columns tracked by ones in $[\partial_d]$ are checked, looking for elements of (h, k) indices with value greater that 2.

```
\langle path\text{-}connected\text{-}cells boundary operator 5a \rangle \equiv
     """ path-connected-cells boundary operator """
     import larlib
     import larcc
     from larcc import *
     def csrBoundaryFilter2(unreliable,out,csrBBMat,cells,FE):
          for row in unreliable:
              for j in range(len(cells)):
                   if out[row,j] == 1:
                        cooCE = csrBBMat.T[j].tocoo()
                       flawedCells = [cooCE.col[k] for k,datum in enumerate(cooCE.data)
                            if datum>2]
                       if all([facet in flawedCells for facet in FE[row]]):
                            out[row, j]=0
          return out
     def csrBoundaryFilter3(unreliable,out,csrBBMat,cells,FE):
          for col in unreliable:
              cooCE = csrBBMat.T[col].tocoo()
              flawedCells = [cooCE.col[k] for k,datum in enumerate(cooCE.data)
                            if datum>2]
              for j in range(out.shape[0]):
                   if out[j,col] == 1:
                       if all([facet in flawedCells for facet in FE[j]]):
                            out[j,col]=0
          return out
Fragment defined by 4, 5a.
Fragment referenced in 29b.
\langle From \ cells \ and \ facets \ to \ boundary \ cells \ 5b \rangle \equiv
```

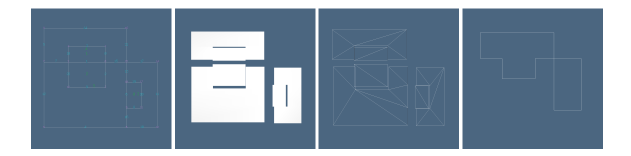


Figure 1: Non-convex LAR 2-complex with (two) 1-cells that are subsets of 2-cells without being their facets. Correctly disentangled by the larUnsignedBoundary2() function: (a) Indexing of 0-, 1-, and 2-cells; (b) exploded 2-cells; (c) triangulated and exploded 2-cells; (d) boundary of the 2-chain [1,1,1,1,0].

```
def totalChain(cells):
   return csr_matrix(len(cells)*[[1]])
def boundaryCells(cells,facets):
    csrBoundaryMat = larBoundary(cells,facets)
   csrChain = csr_matrix(totalChain(cells))
    csrBoundaryChain = csrBoundaryMat * csrChain
   out = [k for k,val in enumerate(csrBoundaryChain.data.tolist()) if val == 1]
   return out
def boundary2Cells(cells,facets,faces):
    csrBoundaryMat = larUnsignedBoundary2(cells,facets,faces)
   csrChain = csr_matrix(totalChain(cells))
   csrBoundaryChain = csrBoundaryMat * csrChain
   out = [k for k,val in enumerate(csrBoundaryChain.data.tolist()) if val == 1]
   return out
def boundary3Cells(cells,facets,faces):
    csrBoundaryMat = boundary3(cells,facets,faces)
   csrChain = csr_matrix(totalChain(cells))
   csrBoundaryChain = csrBoundaryMat * csrChain
   out = [k for k,val in enumerate(csrBoundaryChain.data.tolist()) if val == 1]
   return out
```

Fragment referenced in 29b.

2.3 Correctness proof

Our goal is to get a constructive and, of course, correct representation of the matrix $[\partial_3]$ starting only from M_1 , M_2 , and M_3 .

We have sufficient evidence here to support the correctness of our identification of the matrices of boundary operators as discussed in the previous section. Remember that M_1 , M_2 , and M_3 are the characteristic matrices of 1-cells, 2-cells and 3-cells as subsets of vertices, and that C_0 , C_1 , C_2 , C_3 are the linear spaces of 0-, 1-, 2-, and 3-chains, with coefficients in the field $\mathbb{Z}_2 = \{0,1\}$.

In the following we give a dimension-independent proof, even our implementation is currently restricted to $d \in \{1, 2, 3\}$.

Preamble It is well known that a linear transformation $T: V \to W$ between two linear spaces, with fixed bases of dimension n and m respectively, is represented uniquely by a matrix $A \in \mathbb{R}^{m \times n}$, that by columns contains the coordinate representations in W of the basis vectors of V.

Theorem 1. Consider the linear transformation $\partial_d : C_d \to C_{d-1}$. Having fixed the bases, made by singleton cells in C_d and C_{d-1} , the transformation ∂_d can be represented as a matrix product $c \mapsto [\partial_d] c$, where $c \in C_d$ is the coordinate representation of a d-chain, and $[\partial_d]$ is the $m \times n$ binary matrix having by columns the coordinate representation in C_{d-1} of the boundary (d-1)-chains of cells in C_d .

Proof. The computation of the boundary operator matrix $A_d = [\partial_d] \in \mathbb{Z}_2^{m \times n}$, where m and n are the dimensions of the linear spaces C_d and C_{d-1} , respectively, is made in two steps, in the general case.

First step Consider the characteristic matrices $M_d \in \mathbb{Z}_2^{m \times q}$ and $M_{d-1} \in \mathbb{Z}_2^{n \times q}$ having as rows the images of characteristic functions of bases elements as subsets of vertices, where q is the number of vertices.

Then consider the product matrix $M = M_{d-1}M_d^t = (m_{ij})$, with values in the set \mathbb{N} of non-negative integers. Clearly m_{ij} will coincide with the number of vertices shared by cells $c_i \in C_{d-1}$ and $c_j \in C_d$, i.e. with the cardinality of their intersection as discrete sets of vertices. The predicate

$$m_{ij} \equiv |c_i| \tag{1}$$

is a necessary condition for c_i to be a facet of c_j . Using the Iverson bracket notation, where [P] returns either 1 or 0 depending on the truth of predicate P, we can assign (to) the (i,j) element of our (tentative) boundary matrix the corresponding value:

$$A_d(i,j) := [m_{ij} \equiv |c_i|], \qquad 1 \le i \le m, \ 1 \le j \le n.$$
 (2)

Unfortunately, condition (1) is also *sufficient* for c_i being a facet of c_j only when both are convex cells. In other words, Equation (2) with $A_d = [\partial_d]$ holds in full generality if and

only if the cellular complex under consideration is made only by convex cells. With more general cells, the column j of the ∂_d matrix contains the coordinate representation in C_{d-1} of a possibly proper superset of $\partial_d(c_j)$.

Second step In order to reduce the columns of the approximate boundary matrix A_d to their exact value in \mathbb{Z}_2^m , with $m = \dim C_{d-1}$, we may compute an (approximate) matrix $B \in \mathbb{N}^{p \times n}$ of the operator $\partial_{d-1} \circ \partial_d : C_d \to C_{d-2}$, with $p = \dim C_{d-2}$, by using the approximate representation A_d of the matrix $[\partial_d]$. Let us remark that the exact value of the latter is yet unknown at this point.

According to what asserted in the preamble, every column of this matrix should contain the coordinate representation (in C_{d-2}) of the boundary of the boundary of a basis element in C_d , i.e. of a singleton d-chain.

Therefore, we will enforce the validity of the constraints $\partial \partial = 0$, by checking the values of columns in the product matrix $B_d = A_{d-1}A_d \in \mathbb{N}^{p \times n}$. In particular, every unit vector e_j , *i.e.*, the coordinate representation of a singleton chain $c_j \in C_d$, should be mapped by B_d to the zero vector in \mathbb{Z}_2^p :

$$e_j \mapsto B_d \, e_j = 0^p \in \mathbb{Z}_2^p, \qquad c_j \in C_d.$$
 (3)

Now, let consider the cells of LAR both as subsets of vertices and as cells of a cellular complex. Actually, in the LAR general case, there may be some (d-1)-facets of some d-cell that are subsets of other d-cells of which they are not faces. It is not difficult to provide some examples of this fact (see Figure ??).

When considering each column of B as the coordinate representation of the boundary (d-2)-chain of a d-cell, and noting that every d-cell must be orientable, and hence separating an interior space from an exterior space, we conclude that the number of occurrencies of each (d-2)-cell in a B column must be necessarily even. In particular, it must be either equal to 2 if the column (the d-cell) is locally manifold, or equal to some even number > 2 if the d-cell is locally non-manifold. But odd incidencies of (d-2)-cells along the d-cell boundary are not allowed.

Therefore, for each column B_j $(1 \le j \le n)$ we look for the subset of rows (i.e. boundary (d-2)-faces of c_j) of odd value. If there are none, the column A_j , associated to c_j , is a correct representation of $[\partial_d]_j$. Otherways, we must look for the subsets of (d-1)-cells incorrectly considered facets of c_j .

Let us call R_j the subset of row indices corresponding to odd values in column B_j , and consider the subset of A_j rows with value $a_{ij} = 1$, i.e. the superset S_j of (d-1)-cells including the boundary (d-1)-chain of c_j . The redundant vertex subsets that are not boundary facets of c_j are easily discovered by looking at the (d-2)-boundary of each $s \in S_j$.

In particular, the (d-1)-cell $s \in S_j$ is redundant or extraneous with respect to the c_j boundary if and only if $\partial_{d-1}(s) \subseteq R_j$, because that property has certainly introduced a spurious increment for each (d-2)-facet in the count of incidences of boundary facets.

At this point, we can finally compute the actual boundary matrix ∂_d for a general LAR cellular complex, using again the Iverson brackets:

$$[\partial_d]_{ij} = [(a_{ij} = 1) \land ((R_i = \emptyset) \lor (R_i \not\supseteq \partial_{d-1}(s_i)))] \qquad 1 \le i \le m, \ 1 \le j \le n$$

In words it sounds that the element (i,j) of the boundary matrix $[\partial]_d$ equals that of the "approximate" matrix A_d if and only if either the redundant set R_j is empty, or if it does not contain the (d-1)-boundary chain of the i-th (d-1)-cell.

Just remember that the redundant set R_j contains the (d-2)-faces with odd incidencies on $c_j \in C_d$, computed via the "approximate" matrix $B_d = A_{d-1} \circ A_d$.

2.4 Oriented operators

2.4.1 Oriented simplicial complexes

2.4.2 Oriented LAR complexes

3 From relations to operators

The LAR approach to topology, implemented in the LarLib modules, allows the user to consider the topological relations of incidence and adjacency between faces of a cellular complex as *linear operators* between chains of cells of various dimensions.

The previous approach was to consider the incidence and adjaceny as set-theoretical relations, to be solved by using typical database tools. Conversely, according to the novel IT evolution towards big data and cloud-based storage, even for geometrical data, LarLib takes advantage of a conceptual framework based on linear-algebra and sparse matrices.

3.1 Classification of operators

In the standard solid modeling approach, mainly based on boundary representations, the standard topological operations concern the answers to queries, by reporting the subsets of boundary elements which are incident (different dimension) or adjacent (equal dimension) to assigned boundary elements. Boundary elements stand there for three type of boundary cells: aka $faces\ F$, $edges\ E$, and $vertices\ V$. Nine binary relations may be considered, that are summarized in Table 1.

Conversely, LAR models are normally based on cellular 3-complexes, so using four sets of cells, namely 3-cells C, 2-cells F, 1-cells E, and 0-cells V. The resulting tables of topological relations, and the associated linear operators, are shown in Tables 2a and 2b, respectively.

Table 1: Binary topological relations between boundary elements in boundary representations of solid models

	F	E	V
F	FF	FE	FV
E	EF	EE	EV
V	VF	VE	VV

Table 2: Binary topological relations between cells of LAR decompositions of solid models and corresponding topological operators on ∂_{\circ} chains.

	С	F	Ε	V
С	CC	$\overline{\mathrm{CF}}$	CE	CV
F	FC	$\overline{\mathrm{FF}}$	$\overline{\mathrm{FE}}$	FV
E	EC	EF	EE	EV
V	$\overline{ m VC}$	$\overline{ m VF}$	VE	VV

	С	F	Ε	V
С	$1_C^ op \circ 1_C$	∂_3	$\partial_2 \circ \partial_3$	1_C
F	δ_2	$1_F^\top \circ 1_F$	∂_2	1_F
\mathbf{E}	$\delta_2 \circ \delta_1$	δ_1	$1_E^\top \circ 1_E$	1_{E}
V	$1_C^ op$	1_F^\top	$1_E^ op$	1_V

3.2 Topological relations

3.2.1 Adjacency relations

Fragment referenced in 10b.

Adjacency relations examples

"test/py/boundary/test09.py" $10b\equiv$

```
""" Adjacency relations examples """
from larlib import *

sys.path.insert(0, 'test/py/boundary/')
from test07 import *

⟨kfaces-to-kfaces relation 10a⟩
print "\nCC =",CC
print "\nFF =",FF
print "\nEE =",EE,"\n"

V,BF,BE = larUnsignedBoundary3(V,CV,FV,EV)([1,0])
VIEW(STRUCT(MKTRIANGLES((V,[FV[h] for h in FF[-1]],EV),color=True)))
VIEW(STRUCT(MKPOLS((V,[EV[h] for h in EE[-1]]))+[COLOR(RED)(MKPOLS((V,[EV[-1]]))[0])]))
```

3.2.2 Incidence relations

```
⟨ mfaces-to-nfaces relations 11a⟩ ≡

""" mfaces-to-nfaces relations """

fcOp = larCells2Faces(CV,FV,EV)

CF = [fcOp([k]) for k in range(len(CV))]

FC = invertRelation(CF)

ecOp = larCells2Edges(CV,FV,EV)

CE = [ecOp([k]) for k in range(len(CV))]

EC = invertRelation(CE)

efOp = larFaces2Edges(FV,EV)

FE = [efOp([k]) for k in range(len(FV))]

EF = invertRelation(FE)

⋄
```

Incidence relations examples

Fragment referenced in 11b, 17a.

```
"test/py/boundary/test10.py" 11b≡
""" Incidence relations examples """
from larlib import *
```

```
sys.path.insert(0, 'test/py/boundary/')
from test08 import *

\( \text{mfaces-to-nfaces relations} \) 11a \( \text{print "\nFC =",FC} \)
print "\nEC =",EC
print "\nEF =",EF,"\n"
\( \text{ord} \)
```

3.3 Querying

The more important topological operations in a geometric system concern the answer to queries of the type: "what is the h-chain whose cells are (k-h)-incident to a given h-chain"?

An efficient answer is given by the three higher-level functions in this section, respectively denoted as larCells2Faces, larCells2Edges, and larFaces2Edges. Their first application, over the two or three necessary (compressed) characteristic matrices, returns the csr_matrix of the topological operator, that can be so cached by the calling code. The second application, over the list of h-chain indices, returns the list of k-chain indices of the cells that share with them a (k-h)-face.

3.3.1 Topological incidences

Query from 3-chain to incident 2-chain

```
⟨ Query from 3-chain to incident 2-chain 12⟩ ≡
""" Query from 3-chain to incident 2-chain """

def larCells2Faces(CV,FV,EV):
    csrFC = boundary3(CV,FV,EV)
    def larCells2Faces0(chain):
        chainCoords = csc_matrix((csrFC.shape[1],1),dtype='b')
        for k in chain: chainCoords[k,0] = 1
        out = csrFC * chainCoords
        return out.tocoo().row.tolist()
    return larCells2Faces0
```

Query from 3-chain to incident 1-chain

```
⟨ Query from 3-chain to incident 1-chain 13a⟩ ≡
""" Query from 3-chain to incident 1-chain """

def larCells2Edges(CV,FV,EV):
    lenV = max(CAT(CV))+1
    VV = AA(LIST)(range(lenV))
    csrEC = larUnsignedBoundary2(FV,EV,VV) * boundary3(CV,FV,EV)
    def larCells2Faces0(chain):
        chainCoords = csc_matrix((csrEC.shape[1],1),dtype='b')
        for k in chain: chainCoords[k,0] = 1
        out = csrEC * chainCoords
        return out.tocoo().row.tolist()
    return larCells2Faces0
```

Fragment referenced in 29b.

Compute the signed 2-boundary matrix Compute the signed 2-boundary matrix. The second definition extends the first one by considering LAR faces including holes. For this purpose the signed areas of all loops inside each lar face in FV are taken into account, and only the greates positive area and all the negative areas (except the most negative one) are considered.

```
\langle Compute the signed 2-boundary matrix 13b \rangle \equiv
     """ Compute the signed 2-boundary matrix """
     import triangulation, integr
     def larSignedBoundary2(V,FV,EV):
         efOp = larFaces2Edges(V,FV,EV)
         FE = [efOp([k]) for k in range(len(FV))]
         data,row,col = [],[],[]
         for f in range(len(FE)):
             Vcycles,Ecycles = triangulation.makeCycles((V,[EV[e] for e in FE[f]]))
             Ecycles = [[FE[f][e] for e in cycle] for cycle in Ecycles]
             #areas = integr.signedSurfIntegration((V,Vcycles,EV),signed=True)
             areas = integr.signedSurfIntegration((V, Vcycles, EV), False)
             sortedAreas = sorted((area,k) for k, area in enumerate(areas))
             innerLoops = [zip(Vcycles[k], Ecycles[k]) for area,k in sortedAreas[1:] if area<0]
             outerLoop = [zip(Vcycles[sortedAreas[-1][1]], Ecycles[sortedAreas[-1][1]])]
             orientedFaceLoops = CAT(outerLoop+innerLoops)
             coefficients = [1 if v==EV[e][0] else -1 for v,e in orientedFaceLoops]
```

```
ecycle = [e for v,e in orientedFaceLoops]
  data += coefficients
  row += ecycle
  col += [f]*len(ecycle)
    #print f,len(data),len(row),len(col)
  signedBoundary2 = coo_matrix((data,(row,col)), shape=(len(EV),len(FV)),dtype='b')
  return csr_matrix(signedBoundary2)
```

Fragment referenced in 29b.

Compute any signed 1-boundary chain The script below computes the signed boundary 1-cycle of any 2-chain in a 2-complex. The script returns the pair of arrays orientations, boundary Cells, providing the signs of 1-cells in the extracted 1-cycle and their edge indices, respectively.

Fragment referenced in 29b.

Testing the extraction of a boundary chain An example for testing the extraction of the boundary 1-cycle of a 2-chain $[f_1, f_2]$ is given below. The images generated by the script are displayed in Figure 2.

```
"test/py/boundary/test14.py" 14b=
    """ Testing the extraction of a boundary chain """
    from larlib import *

lines = svg2lines("test/svg2lines/test.svg")
V,FV,EV,polygons = larFromLines(lines,True)
```

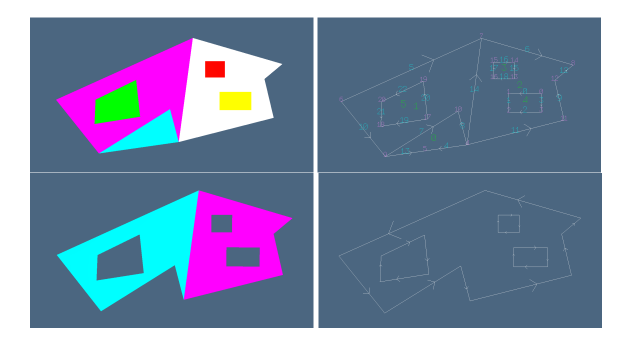


Figure 2: Example of *signed* boundary operator: (a) 2-complex with non-compressible cells; (b) oriented 1-cells; (c) drawing of the 2-chain $[f_1, f_2]$; (d) oriented boundary 1-chain of 2-chain $[f_1, f_2]$.

Query from 2-chain to incident 1-chain

 $\langle Query from 2-chain to incident 1-chain 15 \rangle \equiv$

```
""" Query from 2-chain to incident 1-chain """
def larFaces2Edges(V,FV,EV):
    VV = AA(LIST)(range(len(V)))
    csrEF = larUnsignedBoundary2(FV,EV,VV)
    def larCells2Faces0(chain):
        chainCoords = csc_matrix((csrEF.shape[1],1),dtype='b')
        for k in chain: chainCoords[k,0] = 1
        out = csrEF * chainCoords
        return out.tocoo().row.tolist()
    return larCells2Faces0
```

Fragment referenced in 29b.

3.3.2 Topological adjacencies

kfaces-to-kfaces relations

```
\langle kfaces-to-kfaces \ relations \ 16 \rangle \equiv
     """ kfaces-to-kfaces relations """
     def larCells2Cells(CV,FV,EV):
         csrMat = boundary3(CV,FV,EV)
         csrCC = csrMat.T * csrMat
         def larCells2Cells0(chain):
             chainCoords = csc_matrix((csrCC.shape[1],1),dtype='b')
             for k in chain: chainCoords[k,0] = 1
             out = csrCC * chainCoords
             return out.tocoo().row.tolist()
         return larCells2Cells0
     def larFaces2Faces(FV,EV):
         lenV = max(CAT(FV)) + 1
         VV = AA(LIST)(range(lenV))
         csrMat = larUnsignedBoundary2(FV,EV,VV)
         csrFF = csrMat.T * csrMat
         def larFaces2Faces0(chain):
             chainCoords = csc_matrix((csrFF.shape[1],1),dtype='b')
             for k in chain: chainCoords[k,0] = 1
             out = csrFF * chainCoords
             return out.tocoo().row.tolist()
         return larFaces2Faces0
     def larEdges2Edges(EV,VV):
```

```
lenV = len(VV)
csrMat = larBoundary(EV,VV)
csrEE = csrMat.T * csrMat
def larFaces2Faces0(chain):
    chainCoords = csc_matrix((csrEE.shape[1],1),dtype='b')
    for k in chain: chainCoords[k,0] = 1
    out = csrEE * chainCoords
    return out.tocoo().row.tolist()
return larFaces2Faces0
```

Fragment referenced in 29b.

3.4 Oriented operators

3.4.1 Signed 2-boundary

Testing signed 2-boundary

```
"test/py/boundary/test11.py" 17a\[
""" Testing signed 2-boundary """
from larlib import *

sys.path.insert(0, 'test/py/boundary/')
from test10 import *

\( \lambda m faces-to-n faces relations 11a \rangle
\)
signedBoundary2 = larSignedBoundary2(FV,EV)
\( \lambda \)
```

Testing signed 2-boundary

```
"test/py/boundary/test13.py" 17b\(\text{17b}\)
    """ Testing signed 2-boundary """
    from larlib import *

lines = svg2lines("test/svg2lines/test.svg")
    V,FV,EV,polygons = larFromLines(lines,True)
    VIEW(STRUCT(MKTRIANGLES((V,FV,EV),color=True)))
    submodel = mkSignedEdges((V,EV))
```

```
VV = AA(LIST)(range(len(V)))
VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV],submodel,0.25))
B = larSignedBoundary2(V,FV,EV)
for k in range(B.shape[0]):
    print k,B.todense()[k]

VIEW(STRUCT(MKTRIANGLES((V,FV[1:3],EV),color=True)))
```

Example 2-bondary matrix The example in file test13.py, corresponding to Figure 2 produces the following two-dimensional LAR model:

```
 V = [[0.8627, 0.263], [0.7223, 0.263], [0.7223, 0.1857], [0.8627, 0.1857], [0.5422, 0.0489], [0.3613, 0.0238], [0.0, 0.2296], [0.6044, 0.4933], [1.0, 0.382], [0.1896, 0.0], [0.5037, 0.1896], [0.9614, 0.1522], [0.9207, 0.318], [0.7457, 0.3241], [0.7457, 0.3942], [0.6583, 0.3942], [0.6583, 0.3241], [0.3718, 0.157], [0.1704, 0.1259], [0.3541, 0.3141], [0.1748, 0.2296]] \\ EV = [[0, 1], [2, 1], [3, 2], [0, 3], [4, 5], [6, 7], [7, 8], [9, 10], [4, 10], [11, 12], [6, 9], [4, 11], [12, 8], [9, 5], [4, 7], [13, 14], [15, 14], [16, 15], [13, 16], [17, 18], [17, 19], [20, 18], [19, 20]] \\ \\
```

and the following boundary operator matrix:

$$\partial_2 = \begin{pmatrix} 0 & 0 & -1 & 0 & 1 & 0 \\ 0 & 0 & 1 & 0 & -1 & 0 \\ 0 & 0 & 1 & 0 & -1 & 0 \\ 0 & 0 & 1 & 0 & -1 & 0 \\ 0 & 0 & 1 & 0 & -1 & 0 \\ -1 & 0 & 0 & 0 & 0 & 0 \\ 0 & -1 & 0 & 0 & 0 & 0 \\ 0 & 0 & -1 & 0 & 0 & 0 \\ 0 & 0 & -1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & -1 & 0 & 0 \\ 0 & 0 & 1 & -1 & 0 & 0 \\ 0 & 0 & 1 & -1 & 0 & 0 \\ 0 & 0 & 1 & -1 & 0 & 0 \\ 0 & 0 & 1 & -1 & 0 & 0 \\ 0 & 0 & 1 & -1 & 0 & 0 \\ 0 & 0 & 1 & -1 & 0 & 0 \\ 0 & 0 & 1 & 0 & 0 & -1 \\ 0 & -1 & 0 & 0 & 0 & 1 \\ 0 & -1 & 0 & 0 & 0 & 1 \end{pmatrix}$$

Transformation from chain coordinates to explicit chain data

```
⟨ Transformation from chain coordinates to explicit chain data 19⟩ ≡
    """ Transformation from chain coordinates to explicit chain data """
    def coords2chain(chainCoords):
        coo = coo_matrix(chainCoords)
        return [(e,val) for e,val in zip(coo.row,coo.data)]
```

Fragment referenced in 29b.

3.5 Offset of 2-faces of a 2D complex

In some applications it is necessary to compute the offset of faces of a 2D complex. This may happen for example in architectural applications, where the walls of building layout

must be computed from its wire-frame design. Another important application aims to make more numerically robust the point-in-polygon containment test. In this case the test is driven against a slightly "grown" version of the polygon. The problem solved by the larOffset2D (model) function given below is to "enlarge" each 2-face of a 2-complex in 2D of a (generally small) constant offset. For this purpose, the larOffset2D function operates as follows:

- 1. translate every 1-face towards its cobounday exterior (see Figure 1);
- 2. compute the parametric line equation for the translated pairs of edge vertices;
- 3. compute the intersection points between of adjacent pairs of such lines:
 - (a) solving for both parameters;
 - (b) using the computed parameter value to get the intersection point.

Offset of 2-faces of a 2D complex The coding below is using the Cramer's formula for the the solution of the intersection of two 2D lines via their parametric equation, according to page 50-51 of the formulation given in http://www.ti.inf.ethz.ch/ew/lehre/CG09/materials/v9.pdf. The implementation is simple and very general—e.g.it does not require conditional constructs—so that it may be useful to avoid, for future GPGPU implementations.

```
\langle Offset \ of \ 2-faces \ of \ a \ 2D \ complex \ 20 \rangle \equiv
     """ Offset of 2-faces of a 2D complex """
     from scipy.linalg.basic import det
     def larOffset2D (model,offset=0.001):
         V.FV.EV = model
         newVertices,lines = [],[]
         for f in range(len(FV)):
              # pair of arrays (signs, edges) of f face
              orientations,boundaryCells = larSignedBoundary2Cells(V,FV,EV)([f])
              # array of pairs (sign, edge) of f face
              edges = zip(orientations,boundaryCells)
              # array of pairs (begin_vertex, end_vertex) of oriented edges of f face
              orientedEdges = [tuple(EV[e]) if sign==1 else tuple(REVERSE(EV[e]))
                  for sign, e in edges]
              # array of unit tangentVectors of f face
              tangentVectors = [UNITVECT(VECTDIFF([ V[edge[1]],V[edge[0]] ]))
                  for edge in orientedEdges]
              # array of unit normalVectors of
                                                   f face
             normalVectors = [ SCALARVECTPROD([ offset,[vect[1],-vect[0]] ])
                  for vect in tangentVectors]
```

```
# array of pairs of moved vertices of oriented edges of f face
    movedEdgesOffLine = [[VECTSUM([V[v],n]) for v in orientedEdges[k]]
        for k,n in enumerate(normalVectors)]
    # successor map ( succ[first] := second ) for verts of oriented edges of $f$
    succ = dict(orientedEdges)
    # dictionary of numerals of oriented edges of $f$ (key = pair of vertices)
    edgeDict = dict([(edge,k) for k,edge in enumerate(orientedEdges)])
    # array of pairs of numerals of intersecting edges
    intersections = [[ edgeDict[(u,v)], edgeDict[(v,succ[v])] ]
        for k,(u,v) in enumerate(orientedEdges)]
    # coupling of data points of intersecting pairs
    linepairs = [[movedEdgesOffLine[11],movedEdgesOffLine[12]]
        for 11,12 in intersections]
    # prepare data for line pairs
    linedata = [[ax,ay,bx,by,cx,cy,dx,dy]
        for [[(ax,ay),(bx,by)],[(cx,cy),(dx,dy)]] in linepairs]
    # assemble intersection determinants
    determinants = [ det(mat([[ax-bx,dx-cx], [ay-by,dy-cy]]))
        for [ax,ay,bx,by,cx,cy,dx,dy] in linedata]
    # parameter pairs by Cramer's rule (for oriented edges of f face)
    alpha = [det(mat([[dx-bx,dx-cx],[dy-by,dy-cy]]))/D if abs(D)>.00001 else 0
        for D,(ax,ay,bx,by,cx,cy,dx,dy) in zip(determinants,linedata)]
    # intersection points
    newvert = [(a*mat(p1)+(1-a)*mat(p2)).tolist()[0]
        for a,[[p1,p2],[q1,q2]] in zip(alpha,linepairs)]
    newedges = [[newvert[u],newvert[v]] for u,v in intersections]
    newVertices += [newvert]
    lines += newedges
return lines
```

Fragment referenced in 29b.

Example of Offset generation for the 2-faces of a 2D complex The result of execution of the test program below is given in Figure 3

```
"test/py/boundary/test15.py" 21\(\text{2-faces of a 2D complex """} from larlib import * lines = svg2lines("test/svg2lines/test.svg")
V,FV,EV,polygons = larFromLines(lines,True)
VIEW(STRUCT(MKTRIANGLES((V,FV,EV),color=True)))
```

```
submodel = mkSignedEdges((V,EV))
VV = AA(LIST)(range(len(V)))
VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV],submodel,0.25))
newEdges = larOffset2D((V,FV,EV),offset=0.01)
VIEW(STRUCT(MKPOLS((V,EV)) + AA(COLOR(YELLOW))(AA(POLYLINE)(newEdges))))
```

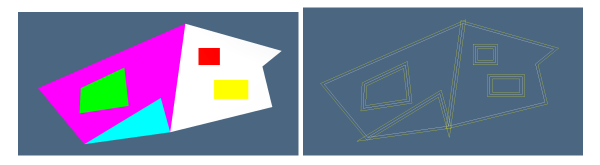


Figure 3: (a) the input 2-complex with general LAR cells; (b) drawing of the offset edges (in yellow) generated by larOffset2D.

3.6 Extraction of 3-cells from a 2-skeleton embedded in 3D

Need to characterize the two vertices of zero edge as "last" of previous edge in the chain, and "first" of following edge in the chain. Therefore the zero edge is oriented as from "last" to "first" and consequently, is "positive" iff "first" > "last".

It is sufficient to characterize the "first" vertex of the edge following the zero edge. This one is the "last" of zero edge. the other vertex of zero edge is its "first". Zero edge is "positive" iff last(zero) > first(zero).

Choose the next face on ordered coboundary of edge Choose the "next" face g_i on "ordered" coboundary of edge

```
⟨ Choose the next face on ordered coboundary of edge 22⟩ =

""" Choose the next face on ordered coboundary of edge """

def adjFace(boundaryOperator,EV,EF_angle,faceChainOrientation):
    def adjFaceO(edge,orientation):
        if orientation > 0: edgeLoop = REVERSE(EF_angle[edge])
```

```
elif orientation < 0: edgeLoop = EF_angle[edge]
edgeLoop = edgeLoop + [edgeLoop[0]] # all positive indices

candidates = set([f for f,_ in faceChainOrientation]).intersection(edgeLoop)
if candidates != set([]):
    pivotFace = candidates.pop()
    if pivotFace in edgeLoop:
        pivotIndex = edgeLoop.index(pivotFace)
    else:
        pivotIndex = edgeLoop.index(-pivotFace)
    adjacentFace = edgeLoop[pivotIndex+1]
else: return None

theSign = boundaryOperator[edge,adjacentFace]
    return adjacentFace, -(theSign*orientation)
return adjFace0
</pre>
```

Choose the start facet in extracting the facet representation of a cell

 \langle Choose the start facet in extracting the facet representation of a cell 23 \rangle \equiv

Fragment referenced in 29b.

```
""" Choose the start facet in extracting the facet representation of a cell """
def chooseStartFace(FV,faceCounter):
    if faceCounter[0,0]==0: return (0,1)
    for f in range(len(FV)):
        if faceCounter[f,0] == 1 and faceCounter[f,1] == 0: return (f,-1)
        elif faceCounter[f,0] == 0 and faceCounter[f,1] == 1: return (f,1)
    for f in range(len(FV)):
        if faceCounter[f,0] == 0 and faceCounter[f,1] == 0: return (f,1)
    if sum(array(faceCounter)) == 2*len(FV): return (-1,999)
    else: print "ERROR: chooseStartFace"
def chooseStartFace(FV,faceCounter):
    for f in range(len(FV)):
        if faceCounter[f,0]==1 and faceCounter[f,1]==0: return (f,-1)
        elif faceCounter[f,0] == 0 and faceCounter[f,1] == 1: return (f,1)
    for f in range(len(FV)):
        if faceCounter[f,0] == 0 and faceCounter[f,1] == 0: return (f,1)
    if sum(array(faceCounter))==2*len(FV): return (-1,999)
    else: return (0,1)
```

Extract the signed representation of a basis element

Fragment referenced in 29b.

Algoritm The algorithm to compute the signed boundary matrix of a 3-complex is given below in pseudocode.

Return the signed boundary matrix of a 3-complex

Algorithm 1 Compute the signed ∂_3 matrix of a 3-complex, from 2-skeleton

```
1: function LARSIGNEDBOUNDARY3(LarModel)
         V, FV, EV \leftarrow \text{LarModel}
 2:
         [\partial_2] \equiv [\delta_1]^t \leftarrow \text{LARSIGNEDBOUNDARY2}(\text{LarModel})
 3:
         sort: EV \to FV \to \mathbb{R}^*: e \mapsto \delta_1(e) (ordered loops of signed 2-faces)
 4:
         [\partial_3] \equiv [\delta_2]^t \leftarrow [] (boundary 2-faces of 3-cells by row)
 5:
         S = \text{indices of 2-faces}
 6:
         while S \neq \emptyset (set of non-traversed 2-faces) do
 7:
              f \leftarrow \text{CHOOSE}(S) (first 2-face, reversing previous sign)
 8:
              F \leftarrow \{f\} (singleton 2-chain)
 9:
              while \partial_2(F) \neq \emptyset do (F non closed)
10:
                   E \leftarrow \partial_2(F) (1-cycle of signed edges)
11:
                   while E \neq \emptyset do
12:
                        for each e \in E do
13:
                            f_i \leftarrow next(f)(sort(\delta_1(e)))
14:
                            F \leftarrow F \cup \{f_i\}
15:
                            S \leftarrow S - \{f_i\} (non-traversed 2-faces)
16:
                            E \leftarrow E - \{e\}
17:
                        end for
18:
                        E \leftarrow \partial_2(F) (new oriented 1-cycle)
19:
                   end while
20:
              end while
21:
              [\partial_3] \leftarrow [\partial_3] + [F] (put F in new [\partial_3] column, i.e. new [\delta_2] row)
22:
         end while
23:
24:
         return [\partial_3] (operator's matrix)
25: end function
```

```
while True:
    startFace,orientation = chooseStartFace(FV,faceCounter)
    if startFace == -1: break
    nonWorkedFaces = nonWorkedFaces.difference({startFace})
    faceChainOrientation = {(startFace, orientation)}
    vect = csc_matrix((m,1),dtype='b')
    for face, orientation in faceChainOrientation:
        vect[face] = orientation
    edgeCycleCoords = boundaryOperator * vect
    edgeCycle = coords2chain(edgeCycleCoords)
    while edgeCycle != []:
        look4face = adjFace(boundaryOperator,EV,EF_angle,faceChainOrientation)
        for edge,orientation in edgeCycle:
            outPair = look4face(edge,orientation)
            if outPair != None:
                adjacentFace, orientation = outPair
                faceChainOrientation = faceChainOrientation.union(
                    [(adjacentFace, orientation)])
                nonWorkedFaces = nonWorkedFaces.difference([adjacentFace])
        vect = csc_matrix((m,1),dtype='b')
        for face, orientation in faceChainOrientation:
            vect[face] = orientation
        edgeCycleCoords = boundaryOperator * vect
        edgeCycle = coords2chain(edgeCycleCoords)
        #if edgeCycle!=[]: VIEW(STRUCT(MKPOLS((V,[EV[e] for e in TRANS(edgeCycle)[0]]))))
    for face, orientation in faceChainOrientation:
        if orientation == 1: faceCounter[face,0]+=1
        elif orientation == -1: faceCounter[face,1]+=1
    #VIEW(STRUCT(MKPOLS((V,[FV[f] for f in TRANS(faceChainOrientation)[0]]))))
    lastrow = [face for face,_ in faceChainOrientation]
    lastcol = [cellNumber for face, orientation in faceChainOrientation]
    lastdata = [orientation for _,orientation in faceChainOrientation]
    lastlength = len(lastrow)
    if lastlength >= longestLength:
        lastrow,longestrow = longestrow,lastrow
        lastcol,longestcol = longestcol,lastcol
        lastdata,longestdata = longestdata,lastdata
        lastlength,longestLength = longestLength,lastlength
    if lastlength != 0:
        row += lastrow
        col += lastcol
        data += lastdata
        CF += [lastrow]
        cellNumber += 1
```

Return the signed boundary matrix of a 3-complex

```
\langle Return \ the \ signed \ boundary \ matrix \ of \ a \ 3-complex \ 27 \rangle \equiv
     """ Return the signed boundary matrix of a 3-complex """
     import boolean
     def larSignedBoundary3((V,FV,EV)):
         model = V, FV, EV
         faceCounter = zeros((len(FV),2),dtype='b')
         CF,m = [],len(FV)
         efOp = larFaces2Edges(V,FV,EV)
         FE = [efOp([k]) for k in range(len(FV))]
         EF_angle, _,_, = boolean.faceSlopeOrdering(model,FE)
         nonWorkedFaces,coboundary_2,cellNumber = set(range(m)),[],0
         boundaryOperator = larSignedBoundary2(V,FV,EV)
         FEbasis = signedBasis(boundaryOperator)
         row,col,data = [],[],[]
         while True:
             startFace,orientation = chooseStartFace(FV,faceCounter)
             if startFace == -1: break
             nonWorkedFaces = nonWorkedFaces.difference({startFace})
             faceChainOrientation = {(startFace, orientation)}
             vect = csc_matrix((m,1),dtype='b')
             for face,orientation in faceChainOrientation:
                  vect[face] = orientation
             edgeCycleCoords = boundaryOperator * vect
              edgeCycle = coords2chain(edgeCycleCoords)
             while edgeCycle != []:
                  look4face = adjFace(boundaryOperator,EV,EF_angle,faceChainOrientation)
                  for edge, orientation in edgeCycle:
                      outPair = look4face(edge,orientation)
                      if outPair != None:
                          adjacentFace, orientation = outPair
                          faceChainOrientation = faceChainOrientation.union(
                              [(adjacentFace, orientation)])
                          nonWorkedFaces = nonWorkedFaces.difference([adjacentFace])
```

```
vect = csc_matrix((m,1),dtype='b')
                 for face, orientation in faceChainOrientation:
                     vect[face] = orientation
                 edgeCycleCoords = boundaryOperator * vect
                 edgeCycle = coords2chain(edgeCycleCoords)
                 #if edgeCycle!=[]: VIEW(STRUCT(MKPOLS((V,[EV[e] for e in TRANS(edgeCycle)[0]]))))
             row += [face for face,_ in faceChainOrientation]
             col += [cellNumber for face,orientation in faceChainOrientation]
             data += [orientation for _,orientation in faceChainOrientation]
             cellNumber += 1
             for face, orientation in faceChainOrientation:
               if orientation == 1: faceCounter[face,0]+=1
               elif orientation == -1: faceCounter[face,1]+=1
             print "faceChainOrientation =",faceChainOrientation
             print "faceCounter =",faceCounter
             CF += [[face for face,_ in faceChainOrientation]]
             print "CF =",CF
             print "\nfaceCounter =",faceCounter
         outMatrix = coo_matrix((data, (row,col)), shape=(m,cellNumber),dtype='b')
         return csr_matrix(outMatrix),CF,faceCounter
     \Diamond
Fragment defined by 24b, 27.
```

Test the signed boundary matrix of a 3-complex

```
\( Test the signed boundary matrix of a 3-complex 28 \) \( = \)
\( """ Test the signed boundary matrix of a 3-complex """ \)
if \( __name__=="__main__": \)
\( V,[VV,EV,FV,CV] = larCuboids([2,2,1],True) \)
\( cubeGrid = Struct([(V,FV,EV)],"cubeGrid") \)
\( cubeGrids = Struct(2*[cubeGrid,t(.5,.5,.5),r(0,0,PI/6)]) \)
\( V,FV,EV = struct2Marshal(cubeGrids) \)
\( VIEW(EXPLODE(1.2,1.2,1.2)(BREP((V,FV,EV),color=False) )) \)
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Fragment referenced in 29b.

Fragment referenced in 29b.

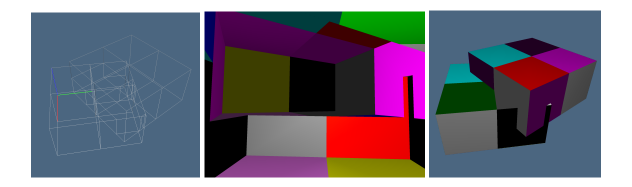


Figure 4: Arrangment of 3-complexes: (a) 1-skeleton of the fragmented (union) 3-complex; (b) view from the interior of the boundary 2-complex; (c) view from the exterior of the 3-complex. Notice that the colour patches correspond to the chain of 2-faces of the boundary of 3-complex.

3.7 Examples

A 3-cell with several holes

4 Exporting

```
"larlib/larlib/boundary.py" 29b≡
""" boundary operators """
from larlib import *
⟨ convex-cells boundary operator 3 ⟩
```

```
\langle path\text{-}connected\text{-}cells boundary operator 4, \dots \rangle
 From cells and facets to boundary cells 5b >
 Marshalling a structure to a LAR cellular model 39 \
 Compute the signed 2-boundary matrix 13b \
 Compute any signed 1-boundary chain 14a
 Offset of 2-faces of a 2D complex 20 \
 Boundary of a 3-complex 40
 Query from 3-chain to incident 2-chain 12 \rangle
 Query from 3-chain to incident 1-chain 13a \
 Query from 2-chain to incident 1-chain 15
 kfaces-to-kfaces relations 16 \
 Transformation from chain coordinates to explicit chain data 19
 Choose the next face on ordered coboundary of edge 22 \
 Choose the start facet in extracting the facet representation of a cell 23 \
 Extract the signed representation of a basis element 24a \
\langle Return \ the \ signed \ boundary \ matrix \ of \ a \ 3-complex \ 24b, \dots \rangle
⟨ Test the signed boundary matrix of a 3-complex 28⟩
```

5 Testing

5.1 Non-oriented operators

Correct boundary extraction example The larBoundary() operator is applied here to a cellular 2-complex of convex cells, producing correct result. It is worth noting that the operator is dimension-independent, and must be applied to the *pair* of compressed characteristic matrices M_d and M_{d-1} , that — in list format — we call either CV,FV or FV,EV, depending on the dimension (either 3 or 2) of the embedding space.

```
"test/py/boundary/test01.py" 30=
    """ testing boundary operators (correct result) """
    from larlib import *

filename = "test/svg/inters/boundarytest0.svg"
    lines = svg2lines(filename)
    VIEW(STRUCT(AA(POLYLINE)(lines)))

V,FV,EV,polygons = larFromLines(lines)
    VV = AA(LIST)(range(len(V)))
    submodel = STRUCT(MKPOLS((V,EV)))
    VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV],submodel,0.2))
    VIEW(EXPLODE(1.2,1.2,1.2)(MKPOLS((V,[EV[e] for e in boundaryCells(FV,EV)],))))
```

```
VIEW(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,FV,EV))))
boundaryOp = larUnsignedBoundary2(FV,EV,VV)

for k in range(1,len(FV)+1):
    faceChain = k*[1]
    BF = chain2BoundaryChain(boundaryOp)(faceChain)
    VIEW(STRUCT(MKPOLS((V,[EV[e] for e in BF]))))
```

Wrong boundary extraction example The larBoundary() operator, applied to a cellular 2-complex wih some non-convex cells, produces incorrect results. In such cases a correct result may be produced only by chance (sometimes this happens). So, be careful to use it only when the precondition (of cell convexity) is everywhere verified. In order to get always a correct result, use the larUnsignedBoundary2 operator.

```
"test/py/boundary/test02.py" 31=
     """ testing boundary operators (wrong result) """
     from larlib import *
     filename = "test/svg/inters/boundarytest3.svg" # KO (MKTRIANGLES) with boundarytest3 !!!
     #filename = "test/svg/inters/boundarytest4.svg"
     lines = svg2lines(filename)
     VIEW(STRUCT(AA(POLYLINE)(lines)))
     V,FV,EV,polygons = larFromLines(lines)
     VV = AA(LIST)(range(len(V)))
     submodel = STRUCT(MKPOLS((V,EV)))
     VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV],submodel,0.2))
     boundaryOp = larUnsignedBoundary2(FV,EV,VV) # <<===== NB
     #boundaryOp = larBoundary(FV,EV) # <<===== NB</pre>
     BF = chain2BoundaryChain(boundaryOp)([1]*len(FV))
     VIEW(EXPLODE(1.2,1.2,1.2)(MKPOLS((V,[EV[e] for e in BF]))))
     VIEW(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,FV,EV),color=True)))
     VIEW(SKEL_1(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,FV,EV)))))
     11 11 11
     for k in range(1,len(FV)+1):
         faceChain = k*[1]
         boundaryChain = chain2BoundaryChain(boundaryOp)(faceChain)
         VIEW(STRUCT(MKPOLS((V, [EV[e] for e in boundaryChain]))))
```

""" \$

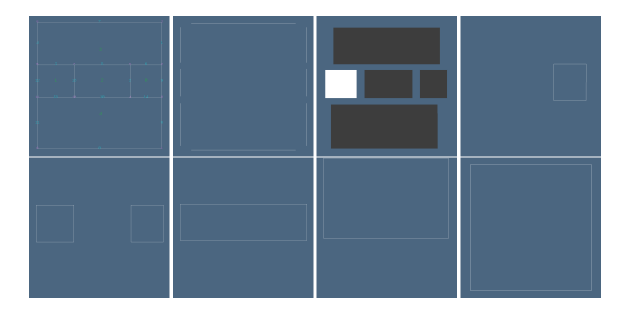


Figure 5: Convex-cell 2-complex. (a) Indexing of 0-,1-,and 2-cells; (b) exploded 2-boundary cells; (c) exploded 2-cells; (d) boundary of a singleton 2-chain; (e-h) boundaries of some 2-chains.

Example Comparison of two implementations of the ∂ operator. Notice the difference between the penultimate rows. In particular, the penultimate row of the matrix generated by larBoundary(FV,EV) is plain wrong. It means that the edge e_{10} is shared by all the (three) 2-cells of the complex. Conversely, it is well known that, for a solid complex, i.e. a d-complex embedded in \mathbb{E}^d , every (d-1)-facet may be shared by no more than 2 d-cells. The resulting boundary of the total chain $[f_0, f_1, f_2]$ codified in coordinates as [1, 1, 1], and shown in Figure 6d, is sonsequently incorrect.

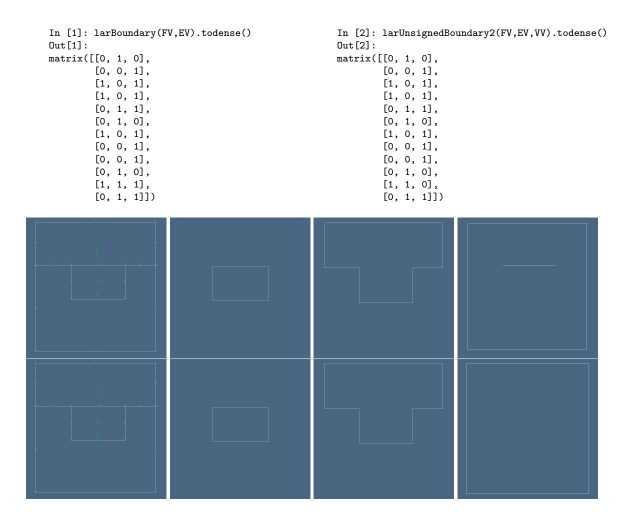


Figure 6: Non-working (i.e. *wrong*) example with boundary. (a) Indexing of 0-,1-,and 2-cells; (b) boundary of a singleton 2-chain; (c) exploded 2-cells; (d) boundary of a singleton 2-chain. Working (i.e. *exact*) example using larUnsignedBoundary2: (e-h) as above.

3D non-convex LAR cells In this example and in the next one we show the boundary computation of LAR models with non-contractible 3- and 2-cells.

```
"test/py/boundary/test03.py" 33 =
    """ 3D non-convex LAR cells """
    from larlib import *

V = [[0.25,0.25,0.0],[0.25,0.75,0.0],[0.75,0.75,0.0],[0.75,0.25,0.0],[1.0, 0.0,0.0],
    [0.0,0.0,0.0],[1.0,1.0,0.0],[0.0,1.0,0.0],[0.25,0.25,1.0],[0.25, 0.25,2.0],[0.25,0.75,
```

```
[0.25, 0.75, 1.0], [0.25, 0.75, -1.0], [0.25, 0.25, 0.25, -1.0], [0.75, 0.75, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0], [0.75, 0.25, -1.0],
-1.0], [0.75, 0.25, 1.0], [0.75, 0.75, 1.0], [1.0, 0.0, 1.0], [0.0, 0.0, 1.0], [1.0, 1.0, 1.0],
[0.0,1.0,1.0],[0.75,0.75,2.0],[0.75,0.25,2.0]]
CV = [(0,1,2,3,4,5,6,7,8,11,16,17,18,19,20,21), (0,1,2,3,8,11,16,17),
(0,1,2,3,12,13,14,15), (8,9,10,11,16,17,22,23)
FV = [(2,3,16,17),(6,7,20,21),(12,13,14,15),(0,1,8,11),(1,2,11,17),(0,1,12,13),
(4,6,18,20),(5,7,19,21),(0,3,13,15),(0,3,8,16),(0,1,2,3),
(10,11,17,22),(2,3,14,15),(8,9,16,23),(8,11,16,17),
(1,2,12,14),(16,17,22,23),(4,5,18,19),(8,9,10,11),(
9,10,22,23),(0,1,2,3,4,5,6,7),(8,11,16,17,18,19,20,21)
EV = [(3,15),(7,21),(10,11),(4,18),(12,13),(5,19),(8,9),(18,19),(22,23),(0,3),(1,11),
(16,17),(0,8),(6,7),(20,21),(3,16),(10,22),(18,20),(19,21),(1,2),(12,14),(4,5),(12,14),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,12),(14,1
8,11), (13,15), (16,23), (14,15), (11,17), (17,22), (2,14), (2,17), (0,1), (9,10), (8,16),
(4,6),(1,12),(5,7),(0,13),(9,23),(6,20),(2,3)
VV = AA(LIST)(range(len(V)))
hpc = STRUCT(MKPOLS((V,EV)))
VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV,CV],hpc,0.6))
BF = boundary3Cells(CV,FV,EV)
VIEW(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,[FV[f]] for f in BF],EV),color=True)))
```

3D non-convex LAR cells In this example the 3D model is constructed partly in automated way, partly by hand. In particular, first we generate a structure of cuboidal complexes, then we transform it is a single complex using part of the computational pipeline being developed for the Boolean arrangments of complexes, so that all the included cells are mutually fragmented. Then the 3-cells are assembed as sets of 2-faces, giving the CF (cells-by-faces) variable. Finally this one is transformed automatically into CV (cells-by-vertices).

```
"test/py/boundary/test04.py" 34\(\text{34}\) """ 3D non-convex LAR cells """ from larlib import * \(\langle Input of a cellular 3-complex 36a \rangle \(\langle Visualization of a 2-chain of a 3-complex 36b \rangle \(\langle Visualization of a 3-chain of a 3-complex 37a \rangle \(\langle \)
```

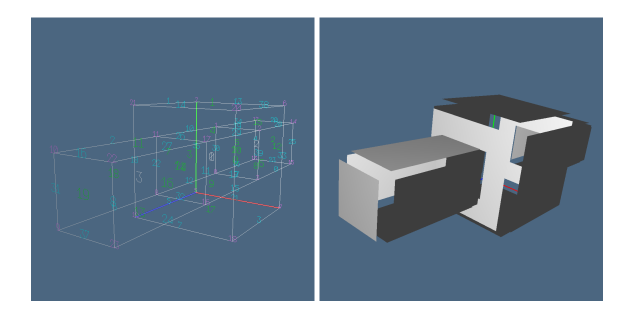


Figure 7: Non-convex 3-complex. (a) Indexing of 0-,1-,2- and 3-cells; (b) exploded 2-boundary cells. Notice that two faces are multiply-connected.

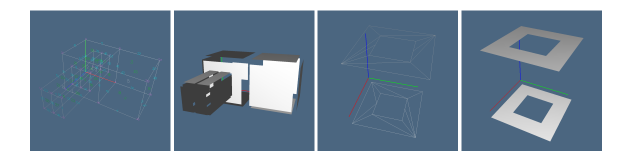


Figure 8: Non-convex 3-complex. (a) Indexing of 0-,1-,2- and 3-cells; (b) exploded 2-boundary cells —notice a drawing error on the back of the model—conversely, the data structures involved are correct, as shown by the two following pictures; (c) solid drawing of the 2-chain [FV[29],FV[30]]; (d) triangulation of the same 2-chain.

Input of a cellular 3-complex

```
⟨Input of a cellular 3-complex 36a⟩ ≡
    """ Input of a cellular 3-complex """
    V,[VV,EV,FV,CV] = larCuboids([2,1,1],True)
    struct = Struct([(V,FV,EV),t(.25,.25,0),s(.25,.5,2),(V,FV,EV)])

V,FV,EV = struct2Marshal(struct)
    CF = AA(sorted)([[20,12,21,5,19,6],[27,1,5,28,13,23],[12,14,25,17,10,4],
    [1,7,17,24,11,18],[30,29,26,16,8,22,10,11,4,18,24,25],[2,3,8,9,0,15]])
    CV = [list(set(CAT([FV[f] for f in faces]))) for faces in CF]

VV = AA(LIST)(range(len(V)))
    hpc = STRUCT(MKPOLS((V,EV)))
    VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV,CV],hpc,0.6))
    ◇
```

Fragment referenced in 34.

Visualization of a 2-chain of a 3-complex

```
\( \text{Visualization of a 2-chain of a 3-complex 36b} \) \( = \)
\( \text{""" Visualization of the boundary 2-chain of a 3-complex """} \)
\( \text{VBF,BE} = \text{larUnsignedBoundary3(V,CV,FV,EV)(len(CV)*[1])} \)
\( \text{VIEW(STRUCT(MKTRIANGLES((V,BF,EV),color=True)))} \)
\( \text{VIEW(SKEL_1(STRUCT(MKTRIANGLES((V,BF,EV)))))} \)
\( \text{boundaryEdges} = \text{chain2BoundaryChain(larUnsignedBoundary2(FV,EV,VV))} \)
\( \text{edgeChain} = \text{boundaryEdges(29*[0]+[1]+[1])} \)
\( \text{VIEW(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,FV[29:31],[EV[e] for e in edgeChain]))))} \)
\( \text{VIEW(SKEL_1(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,FV[29:31],[EV[e] for e in edgeChain])))))} \)
\( \text{"""}
\( \text{"""}
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\( \text{""""}
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\( \text{""""}
```

Fragment referenced in 34.

Visualization of a 3-chain of a 3-complex

```
\langle Visualization of a 3-chain of a 3-complex 37a \rangle \equiv
     """ Visualization of a 3-chain of a 3-complex """
     print "\n***** ECCOMI"
     V,BF,BE = larUnsignedBoundary3(V,CV,FV,EV)([0,0,0,0,1,1])
     VIEW(STRUCT(MKTRIANGLES((V,BF,BE))))
     VIEW(SKEL_1(STRUCT(MKTRIANGLES((V,BF,BE)) )))
Fragment referenced in 34.
"test/py/boundary/test05.py" 37b=
     """ Boundary of a 3-complex """
     from larlib import *
     V,[VV,EV,FV,CV] = larCuboids([1,1,1],True)
     cube = Struct([ (V,FV,EV) ])
     assembly = Struct([ cube, Struct([t(0,.5,0), r(PI/4,0,0), s(.5,.5,.5), cube]) ])
     V,FV,EV = struct2Marshal(assembly)
     VV = AA(LIST)(range(len(V)))
     hpc = STRUCT(MKPOLS((V,EV)))
     VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV],hpc,0.6))
     CF = [[1,2,3,4,6,7],[0,1,2,3,4,5,6,7,8,9,10,11]]
     CV = [list(set(CAT([FV[f] for f in faces]))) for faces in CF]
     V,BF,BE = larUnsignedBoundary3(V,CV,FV,EV)([0,1])
     VIEW(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,BF,BE),color=True)))
"test/py/boundary/test06.py" 37c \equiv
     """ Boundary of a 3-complex """
     from larlib import *
     V,[VV,EV,FV,CV] = larCuboids([1,1,1],True)
     cube = Struct([ (V,FV,EV) ])
     hole = Struct([t(0,.5,0), r(PI/4,0,0), s(.5,.5,.5), cube])
     assembly = Struct([ cube, hole, t(0,0,SQRT(0.5)), hole ])
```

```
V,FV,EV = struct2Marshal(assembly) # WRONG: TODO: check ...
     VV = AA(LIST)(range(len(V)))
     hpc = STRUCT(MKPOLS((V,EV)))
     VIEW(larModelNumbering(1,1,1)(V,[[],[],FV],hpc,0.6))
     CF = [[4,5,7,16,17,19,20],[3,8,6,12,11,13],[0,1,10,20],[]]
     CV = [list(set(CAT([FV[f] for f in faces]))) for faces in CF]
     V,BF,BE = larUnsignedBoundary3(V,CV,FV,EV)([0,1,1,0])
     VIEW(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,BF,BE)))) # ERROR in MKTRIANGLES with non-manifold fa
     VIEW(EXPLODE(1.2,1.2,1.2)(MKFACES((V,BF,EV))))
     VIEW(SKEL_1(EXPLODE(1.2,1.2,1.2)(MKFACES((V,BF,EV)))))
"test/py/boundary/test07.py" 38a=
     """ Boundary of a 3-complex """
     from larlib import *
     V,[VV,EV,FV,CV] = larCuboids([1,1,1],True)
     cube = Struct([ (V,FV,EV) ])
     hole = Struct([t(0,.5,0), r(PI/4,0,0), s(1,.5/SQRT(2),.5/SQRT(2)), cube])
     assembly = Struct([ cube, hole ])
     V,FV,EV = struct2Marshal(assembly) # WRONG: TODO: check ...
     VV = AA(LIST)(range(len(V)))
     hpc = STRUCT(MKPOLS((V,EV)))
     VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV],hpc,0.6))
     CF = [[1,3,6,7,12,11],[0,2,4,5,9,8]]
     CV = [list(set(CAT([FV[f] for f in faces]))) for faces in CF]
     V,BF,BE = larUnsignedBoundary3(V,CV,FV,EV)([1,0])
     VIEW(EXPLODE(1.2,1.2,1.2) (MKTRIANGLES((V,BF,EV),color=True)))
"test/py/boundary/test08.py" 38b\equiv
     """ Boundary of a 3-complex """
```

from larlib import *

```
V, [VV,EV,FV,CV] = larCuboids([1,1,1],True)
cube = Struct([ (V,FV,EV) ])
hole = Struct([t(0,.5,0), r(PI/4,0,0), s(1,.5/SQRT(2),.5/SQRT(2)), cube])
assembly = Struct([ cube, hole ])
assembly2 = Struct([ assembly, t(0,0,.5), s(0.5,1,1), hole ])
V,FV,EV = struct2Marshal(assembly2)
VV = AA(LIST)(range(len(V)))
hpc = STRUCT(MKPOLS((V,EV)))
VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV],hpc,0.7))
\#CF = [[1,3,6,14,17,18,19,0,4,9,11,15,16,2,5,7,8,10,13],[0,4,9,11,15,16],[2,5,7,8,10,13]]
CF = [[16,9,11,0,15,4],[10,12,7,2,17,15,9,5,18,8,16,3,6,14,19,4,1],[5,13,7,10,8,2]]
CV = [list(set(CAT([FV[f] for f in faces]))) for faces in CF]
n = len(CV)
for k in range(n):
   V,BF,BE = larUnsignedBoundary3(V,CV,FV,EV)(IDNT(n)[k])
   VIEW(STRUCT(MKTRIANGLES((V,BF,BE),color=True)))
   VIEW(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,BF,BE),color=True)))
   VIEW(SKEL_1(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,BF,BE)))))
V,BF,BE = larUnsignedBoundary3(V,CV,FV,EV)([1,1,1])
VIEW(SKEL_1(EXPLODE(1.2,1.2,1.2)(MKTRIANGLES((V,BF,BE)))))
```

A Utilities

Marshalling a structure to a LAR cellular model The function struct2Marshal transforms a Struct object, often used to define some assembly of simpler models, to a correctly defined LAR cellular model, i.e. to a cellular partition of the space, in other words a quasi-disjoint partition of the object into well-glued cells of suitable dimensions.

```
⟨ Marshalling a structure to a LAR cellular model 39⟩ ≡

""" Marshalling a structure to a LAR cellular model """

import boolean,inters

def struct2Marshal(struct):
    W,FW,EW = struct2lar(struct)
    quadArray = [[W[v] for v in face] for face in FW]
    parts = boolean.boxBuckets3d(boolean.containmentBoxes(quadArray))
    Z,FZ,EZ = boolean.spacePartition(W,FW,EW, parts)
```

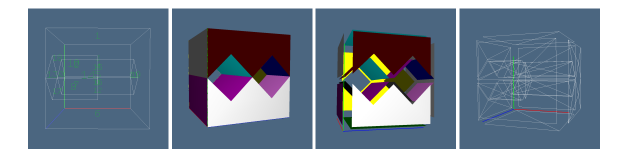


Figure 9: Decomposition of the unit 3-cube in a cellular 3-complex with three 3-cells. Two 3-cells are homeomorphic to the 3-ball, while the remaining one is homeomorphic to the 3-torus. Notice that one of 2-cells (as well one of 3-cells) are non-contractible and non-manifold, as well as non-convex: (a) Indexing of 2-faces of the 3-complex; (b) drawing of the (boundary of) non-convex 3-cell; (c) exploded drawing of the (boundary of) non-convex 3-cell; (d) triangulation of its 2-faces. The triangulation of LAR 2-faces is needed in order to draw them solidly.

```
V,FV,EV = inters.larSimplify((Z,FZ,EZ),radius=0.001)
return V,FV,EV
```

Fragment referenced in 29b.

Boundary of a 3-complex

```
    """ Boundary of a 3-complex """
    import larcc
    """ WHY wrong ???? TOCHECK !!
    def larUnsignedBoundary3(V,CV,FV,EV):
        VV = AA(LIST)(range(len(V)))
        operator3 = larcc.chain2BoundaryChain(boundary3(CV,FV,EV))
        operator2 = larcc.chain2BoundaryChain(larUnsignedBoundary2(FV,EV,VV))
        def larUnsignedBoundary30(chain):
            BF = operator3(chain)
            faceCoords = len(FV)*[0]
            for f in BF: faceCoords[f] = 1
            BE = operator2(faceCoords)
            return V,[FV[f] for f in BF],[EV[e] for e in BE]
        return larUnsignedBoundary30
```

```
def larUnsignedBoundary3(V,CV,FV,EV):
    VV = AA(LIST)(range(len(V)))
    operator3 = larcc.chain2BoundaryChain(boundary3(CV,FV,EV))
    operator2 = larcc.chain2BoundaryChain(larUnsignedBoundary2(FV,EV,VV))
    def larUnsignedBoundary30(chain):
        BF = operator3(chain)
        BE = set()
        for f in BF:
            faceCoords = len(FV)*[0]
            faceCoords[f] = 1
            BE = BE.union(operator2(faceCoords))
        return V,[FV[f] for f in BF],[EV[e] for e in BE]
    return larUnsignedBoundary30
```

Fragment referenced in 29b.

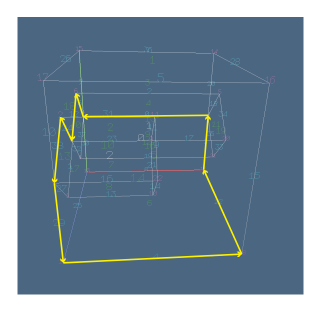


Figure 10: The oriented boundary 1-cycle of a partial 3-cell extraction from a 2-complex in 3D.

References

[CL13] CVD-Lab, $Linear\ algebraic\ representation,$ Tech. Report 13-00, Roma Tre University, October 2013.