

Boolean chains *

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Abstract

A novel algorithm for computation of Boolean operations between cellular complexes is given in this module. It is based on bucketing of possibly interacting geometry using a box-extension of kd-trees, normally used for point proximity queries. Such kd-tree representation of containment boxes of cells, allow us to compute a number of independent buckets of data to be used for local intersection, followed by elimination of duplicated data. Actually we reduce the intersection of boundaries in 3D to the independent intersections of the buckets of (transformed) faces with the 2D subspace $z = 0$, in order to reconstruct each splitted facet of boolean arguments, suitably transformed ther together with the bucket of indent facets. A final tagging of cells as either belonging or not to each operand follows, allowing for fast extraction of Boolean results between any pair of chains (subsets of cells). This Boolean algorithm can be considered of a *Map-Reduce* kind, and hence suitable of a distributed implementation over big datasets. The actual engineered implementation will follow the present prototype, using some distributed NoSQL database, like MongoDB or Riak.

Contents

1	Introduction	2
2	Preview of the algorithm	2
2.1	Unification	2
2.2	Bucketing	2
2.3	Intersection	3
2.4	Reconstruction	3

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3	Implementation	3
3.1	Box-kd-tree	3
3.2	Merging the boundaries	5
3.3	Elementary splitting	5
3.4	Boolean chains	6
4	Esporting the Library	6
5	Test examples	7
5.1	Random triangles	7
5.2	Testing the box-kd-trees	9
5.3	Intersection of geometry subsets	10
A	Code utilities	11

1 Introduction

2 Preview of the algorithm

The whole Boolean algorithm is composed by four stages in sequence, denoted in the following as *Unification*, *Bucketing*, *Intersection*, and *Reconstruction*. The algorithm described here is both multidimensional and variadic. Multidimensional means that the arguments are solid in Euclidean space of dimension d , with d small integer. The *arity* of a function or operation is the number of arguments or operands the function or operation accepts. In computer science, a function accepting a variable number of arguments is called *variadic*.

2.1 Unification

In this first step the boundaries of the n Boolean arguments are computed and merged together as a set of chains defined in the discrete set \mathbf{V} made by the union of their vertices, and possibly by a discrete set of points generated by intersection of cells of complementary dimension, i.e. whose dimensions add up to the dimension of the ambient space. Actually, only the (*oriented*) boundaries $\mathbf{V}, \mathbf{FV}_i$ ($1 \leq i \leq n$) of the various arguments are retained here, and used by the following steps of the algorithm.

2.2 Bucketing

The bounding boxes of facets \mathbf{FV}_i are computed, and their *box-kd-tree* is worked-out, so providing a group of buckets of close cells, that can be elaborated independently, and possibly in parallel, to compute the intersections of the boundary cells.

2.3 Intersection

For each facet f of one of Boolean arguments, the subset $F(f)$ of incident or intersecting facets of boundaries of the other arguments were computed in the previous *bucketing* step. So, each F is transformed by the affine map that sends f into the $z = 0$ subspace, and there is intersected with this subspace, generating a subset $E(f)$ of coplanar edges. This one is projected in 2D, and the *regularized* cellular 2-complex $G(f)$ induced by it is computed, and mapped back to the original space position and orientation of f (providing a partition of it induced by the other boundaries).

2.4 Reconstruction

Like for in the reconstruction of 2D solid cells using the angular ordering of edges around the vertices, the coincident edges are identified in 3D, and used to sort the incident faces using the values of solid angles given with one reference face. The 3D space partition induced by $\cup_f G(f)$ is finally reconstructed, possibly in parallel, by traversing the adjacent sets of facets on the boundary of each solid cell.

3 Implementation

3.1 Box-kd-tree

Split the boxes between the (below,above) subsets

```
⟨ Split the boxes between the below,above subsets 2 ⟩ ≡  
    """ Split the boxes between the below,above subsets """  
    def splitOnThreshold(boxes,subset,coord):  
        theBoxes = [boxes[k] for k in subset]  
        threshold = centroid(theBoxes,coord)  
        ncoords = len(boxes[0])/2  
        a = coord%ncoords  
        b = a+ncoords  
        below,above = [],[]  
        for k in subset:  
            if boxes[k][a] <= threshold: below += [k]  
        for k in subset:  
            if boxes[k][b] >= threshold: above += [k]  
        return below,above  
    ◇
```

Macro referenced in [6b](#).

Test if bucket OK or append to splitting stack

```
⟨ Test if bucket OK or append to splitting stack 3a ⟩ ≡
```

```

""" Test if bucket OK or append to splitting stack """
def splitting(bucket,below,above, finalBuckets,splittingStack):
    if (len(below)<4 and len(above)<4) or len(set(bucket).difference(below))<7 \
        or len(set(bucket).difference(above))<7:
        finalBuckets.append(below)
        finalBuckets.append(above)
    else:
        splittingStack.append(below)
        splittingStack.append(above)

```

◇

Macro referenced in 6b.

Remove subsets from bucket list

⟨ Remove subsets from bucket list 3b ⟩ ≡

```

""" Remove subsets from bucket list """
def removeSubsets(buckets):
    n = len(buckets)
    A = zeros((n,n))
    for i,bucket in enumerate(buckets):
        for j,bucket1 in enumerate(buckets):
            if set(bucket).issubset(set(bucket1)):
                A[i,j] = 1
    B = AA(sum)(A.tolist())
    out = [bucket for i,bucket in enumerate(buckets) if B[i]==1]
    return out

def geomPartitionate(boxes,buckets):
    geomInters = [set() for h in range(len(boxes))]
    for bucket in buckets:
        for k in bucket:
            geomInters[k] = geomInters[k].union(bucket)
    for h,inters in enumerate(geomInters):
        geomInters[h] = geomInters[h].difference([h])
    return AA(list)(geomInters)

```

◇

Macro referenced in 6b.

Iterate the splitting until splittingStack is empty

⟨ Iterate the splitting until splittingStack is empty 4 ⟩ ≡

```

""" Iterate the splitting until \texttt{splittingStack} is empty """
def boxTest(boxes,h,k):
    B1,B2,B3,B4,B5,B6,_ = boxes[k]

```

```

b1,b2,b3,b4,b5,b6,_ = boxes[h]
return not (b4<B1 or B4<b1 or b5<B2 or B5<b2 or b6<B3 or B6<b3)

def boxBuckets(boxes):
    bucket = range(len(boxes))
    splittingStack = [bucket]
    finalBuckets = []
    while splittingStack != []:
        bucket = splittingStack.pop()
        below,above = splitOnThreshold(boxes,bucket,1)
        below1,above1 = splitOnThreshold(boxes,above,2)
        below2,above2 = splitOnThreshold(boxes,below,2)

        below11,above11 = splitOnThreshold(boxes,above1,3)
        below21,above21 = splitOnThreshold(boxes,below1,3)
        below12,above12 = splitOnThreshold(boxes,above2,3)
        below22,above22 = splitOnThreshold(boxes,below2,3)

        splitting(above1,below11,above11, finalBuckets,splittingStack)
        splitting(below1,below21,above21, finalBuckets,splittingStack)
        splitting(above2,below12,above12, finalBuckets,splittingStack)
        splitting(below2,below22,above22, finalBuckets,splittingStack)

        finalBuckets = list(set(AA(tuple)(finalBuckets)))
    parts = geomPartitionate(boxes,finalBuckets)
    parts = [[h for h in part if boxTest(boxes,h,k)] for k,part in enumerate(parts)]
    return AA(sorted)(parts)

```

◇

Macro referenced in 6b.

aaaaaa

```

⟨ aaaaaa 5a ⟩ ≡
    """ aaaaa """

```

◇

Macro never referenced.

3.2 Merging the boundaries

3.3 Elementary splitting

In this section we implement the splitting of $(d-1)$ -faces, stored in **FV**, induced by the buckets of $(d-1)$ -faces, stored in **parts**, and one-to-one associated to them. Of course, (a) both such arrays have the same number of elements, and (b) whereas **FV** contains the

indices of incident vertices for each face, `parts` contains the indices of adjacent faces for each face, with the further constraint that $i \notin \text{parts}(i)$.

Computation of face transformations The faces in every `parts(i)` must be affinely transformed into the subspace $x_d = 0$, in order to compute the intersection of its elements with this subspace, that are submanifolds of dimension $d - 2$.

```

⟨Computation of face transformations 5b⟩ ≡
    """ Computation of affine face transformations """
    def COVECTOR(points):
        pointdim = len(points[0])
        plane = Planef.bestFittingPlane(pointdim,[item for sublist in points for item in sublist])
        return [plane.get(I) for I in range(0,pointdim+1)]

    def faceTransformations(facet):
        covector = COVECTOR(facet)
        translVector = facet[0]
        # translation
        newFacet = [ VECTDIFF([v,translVector]) for v in facet ]
        # linear transformation: boundaryFacet -> standard (d-1)-simplex
        d = len(facet[0])
        transformMat = mat( newFacet[1:d] + [covector[1:]] ).T.I
        # transformation in the subspace x_d = 0
        out = (transformMat * (mat(newFacet).T)).T.tolist()
        print "\nin =",facet
        print "out =",out
        return transformMat

```

◇
Macro referenced in [6b](#).

```

⟨Submanifold maps computation 6a⟩ ≡
    """ Submanifold maps computation """
    for face in FV:
        verts = [V[v] for v in face]
        facet = [ VECTDIFF([v,verts[0]]) for v in verts ]
        transf = faceTransformations(facet)
        new = (transf * (mat(facet).T)).T.tolist()
        VIEW(POLYLINE(new))

```

◇
Macro never referenced.

3.4 Boolean chains

4 Exporting the Library

```

"lib/py/bool2.py" 6b ≡

```

```

""" Module for Boolean computations between geometric objects """
from pyplasm import *
""" import modules from larcc/lib """
import sys
sys.path.insert(0, 'lib/py/')
from inters import *
DEBUG = True

< Coding utilities 11a >
< Split the boxes between the below,above subsets 2 >
< Test if bucket OK or append to splitting stack 3a >
< Remove subsets from bucket list 3b >
< Iterate the splitting until splittingStack is empty 4 >
< Computation of face transformations 5b >
< Computation of affine face transformations ? >
◇

```

5 Test examples

5.1 Random triangles

Generation of random triangles and their boxes

```

"test/py/bool2/test01.py" 6c ≡
    """ Generation of random triangles and their boxes """
    import sys
    sys.path.insert(0, 'lib/py/')
    from bool2 import *
    glass = MATERIAL([1,0,0,0.1, 0,1,0,0.1, 0,0,1,0.1, 0,0,0,0.1, 100])

    randomTriaArray = randomTriangles(10,0.99)
    VIEW(STRUCT(AA(MKPOL)([[verts, [[1,2,3]], None] for verts in randomTriaArray]]))

    boxes = containmentBoxes(randomTriaArray)
    hexas = AA(box2exa)(boxes)
    cyan = COLOR(CYAN)(STRUCT(AA(MKPOL)([[verts, [[1,2,3]], None] for verts in randomTriaArray]]))
    yellow = STRUCT(AA(glass)(AA(MKPOL)([hex for hex,qualifier in hexas]]))
    VIEW(STRUCT([cyan,yellow]))
◇

```

Generation of random quadrilaterals and their boxes

```

"test/py/bool2/test02.py" 7a ≡
    """ Generation of random quadrilaterals and their boxes """
    import sys
    sys.path.insert(0, 'lib/py/')

```

```

from bool2 import *
glass = MATERIAL([1,0,0,0.1, 0,1,0,0.1, 0,0,1,0.1, 0,0,0,0.1, 100])

randomQuadArray = randomQuads(10,1)
VIEW(STRUCT(AA(MKPOL)([[verts, [[1,2,3,4]], None] for verts in randomQuadArray]]))

boxes = containmentBoxes(randomQuadArray)
hexas = AA(box2exa)(boxes)
cyan = COLOR(CYAN)(STRUCT(AA(MKPOL)([[verts, [[1,2,3,4]], None] for verts in randomQuadArray]]))
yellow = STRUCT(AA(glass)(AA(MKPOL)([hex for hex,qualifier in hexas])))
VIEW(STRUCT([cyan,yellow]))
◇

```

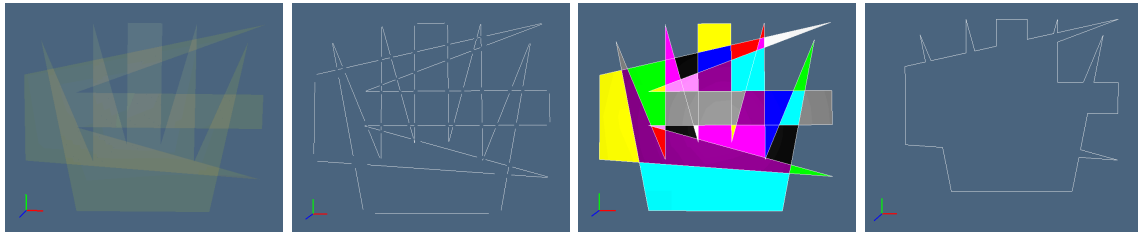


Figure 1: LAR complex from two polygons. (a) the input polygons; (b) the intersection of boundary lines; (c) the extracted *regularized* 2-complex; (d) the boundary LAR.

```

"test/py/bool2/test03.py" 7b ≡
    """ Boolean complex generated by boundaries of two complexes """
    import sys
    sys.path.insert(0, 'lib/py/')
    from inters import *
    glass = MATERIAL([1,0,0,0.1, 0,1,0,0.1, 0,0,1,0.1, 0,0,0,0.1, 100])

    V1 = [[3,0],[11,0],[13,10],[10,11],[8,11],[6,11],[4,11],[1,10],[4,3],[6,4],
           [8,4],[10,3]]
    FV1 = [[0,1,8,9,10,11],[1,2,11],[3,10,11],[4,5,9,10],[6,8,9],[0,7,8]]
    EV1 = [[0,1],[0,7],[0,8],[1,2],[1,11],[2,11],[3,10],[3,11],[4,5],[4,10],[5,
           9],[6,8],[6,9],[7,8],[8,9],[9,10],[10,11]]
    BE1 = boundaryCells(FV1,EV1)
    lines1 = [[V1[v] for v in EV1[edge]] for edge in BE1]

    V2 = [[0,3],[14,2],[14,5],[14,7],[14,11],[0,8],[3,7],[3,5]]
    FV2 = [[0,5,6,7],[0,1,7],[4,5,6],[2,3,6,7]]
    EV2 = [[0,1],[0,5],[0,7],[1,7],[2,3],[2,7],[3,6],[4,5],[4,6],[5,6],[6,7]]
    BE2 = boundaryCells(FV2,EV2)
    lines2 = [[V2[v] for v in EV2[edge]] for edge in BE2]

```



```

VIEW(STRUCT([ glass(STRUCT(MKPOLS((V1,FV1)))), glass(STRUCT(MKPOLS((V2,FV2)))) ]))
lines = lines1 + lines2
VIEW(STRUCT(AA(POLYLINE)(lines)))

global precision
PRECISION += 2
V,FV,EV = larFromLines(lines)
VIEW(EXPLODE(1.2,1.2,1)(MKPOL((V,EV))))

VV = AA(LIST)(range(len(V)))
submodel = STRUCT(MKPOLS((V,EV)))
VIEW(larModelNumbering(1,1,1)(V,[VV,EV,FV[:-1]],submodel,1))

polylines = [[V[v] for v in face+[face[0]]] for face in FV[:-1]]
colors = [CYAN, MAGENTA, WHITE, RED, YELLOW, GREEN, GRAY, ORANGE, BLACK, BLUE, PURPLE, BROWN]
sets = [COLOR(colors[k%12])(FAN(pol)) for k,pol in enumerate(polylines)]
VIEW(STRUCT([ T(3)(0.02)(STRUCT(AA(POLYLINE)(lines))), STRUCT(sets)]))

VIEW(EXPLODE(1.2,1.2,1)((AA(POLYLINE)(polylines))))
polylines = [ [V[v] for v in FV[-1]+[FV[-1][0]]] ]
VIEW(EXPLODE(1.2,1.2,1)((AA(POLYLINE)(polylines))))
◇

```

5.2 Testing the box-kd-trees

Visualizing with different colors the buckets of box-kd-tree

"test/py/bool2/test04.py" 8 ≡

```

""" Visualizing with different colors the buckets of box-kd-tree """
from pyplasm import *
""" import modules from larcc/lib """
import sys
sys.path.insert(0, 'lib/py/')
from bool2 import *

randomQuadArray = randomQuads(30,0.8)
VIEW(STRUCT(AA(MKPOL)([[verts, [[1,2,3,4]], None] for verts in randomQuadArray]]))

boxes = containmentBoxes(randomQuadArray)
hexas = AA(box2exa)(boxes)
glass = MATERIAL([1,0,0,0.1, 0,1,0,0.1, 0,0,1,0.1, 0,0,0,0.1, 100])
yellow = STRUCT(AA(glass)(AA(MKPOL)([hex for hex,data in hexas])))
VIEW(STRUCT([#cyan,
             yellow]))

```

```

parts = boxBuckets(boxes)
for k,part in enumerate(parts):
    bunch = [glass(STRUCT( [MKPOL(hexas[h][0]) for h in part]))]
    bunch += [COLOR(RED)(MKPOL(hexas[k][0]))]
    VIEW(STRUCT(bunch))
◇

```

5.3 Intersection of geometry subsets

Two unit cubes

```

⟨Two unit cubes 9⟩ ≡
    """ Two unit cubes """
    import sys
    sys.path.insert(0, 'lib/py/')
    from bool2 import *
    glass = MATERIAL([1,0,0,0.1, 0,1,0,0.1, 0,0,1,0.1, 0,0,0,0.1, 100])

    V,[VV,EV,FV,CV] = larCuboids([1,1,1],True)
    cube1 = Struct([(V,FV,EV)],"cube1")
    twoCubes = Struct([cube1,t(.5,.5,.5),cube1])
    V,FV,EV = struct2lar(twoCubes)
    VIEW(EXPLODE(1.2,1.2,1.2)(MKPOL((V,FV))))

    quadArray = [[V[v] for v in face] for face in FV]
    boxes = containmentBoxes(quadArray)
    hexas = AA(box2exa)(boxes)
    parts = boxBuckets(boxes)
    ◇

```

Macro referenced in [10](#).

Face (and incident faces) transformation

```

"test/py/bool2/test05.py" 10 ≡
    """ Face (and incident faces) transformation """

    ⟨Two unit cubes 9⟩

    def crossRelation(XV,YV):
        csrXV = csrCreate(XV)
        csrYV = csrCreate(YV)
        csrXY = matrixProduct(csrXV, csrYV.T)
        XY = [None for k in range(len(XV))]
        for k,face in enumerate(XV):
            data = csrXY[k].data

```

```

        col = csrXY[k].indices
        XY[k] = [col[h] for h,val in enumerate(data) if val==2]
    return XY

def mapping(pivotFace):
    tx,ty,tz = pivotFace[0]
    transl = mat([[1,0,0,-tx],[0,1,0,-ty],[0,0,1,-tz],[0,0,0,1]])
    facet = [ VECTDIFF([v,pivotFace[0]]) for v in pivotFace ]
    m = faceTransformations(facet)
    mapping = mat([[m[0,0],m[0,1],m[0,2],0],[m[1,0],m[1,1],m[1,2],0],[m[2,0],m[2,1],m[2,2],0],
    transform = mapping * transl
    return transform

def intersection(V,FV,EV):
    def intersection0(k,bundledFaces):
        FE = crossRelation(FV,EV)
        pivotFace = [V[v] for v in FV[k]]
        transform = mapping(pivotFace) # submanifold transformation
        transformedCells,edges,faces = [],[],[]
        for face in bundledFaces:
            edge = set(FE[k]).intersection(FE[face])
            if edge == set():
                cell = [V[v]+[1.0] for v in FV[face]] # vertices of incident face
                transformedCell = (transform * (mat(cell).T)).T.tolist() # vertices in local
                transformedCells += [[point[:-1] for point in transformedCell]]
                faces = [MKPOL([cell,[range(1,len(cell)+1)],None)] for cell in transformedCell]
            else:
                e, = edge
                vs = [V[v]+[1.0] for v in EV[e]]
                ws = (transform * (mat(vs).T)).T.tolist()
                edges += [POLYLINE([p[:-1] for p in ws])]
        return edges,faces
    return intersection0

for k,bundledFaces in enumerate(parts):
    edges,faces = intersection(V,FV,EV)(k,bundledFaces)
    VIEW(STRUCT(edges + AA(glass)(faces)))

```

◇

A Code utilities

Coding utilities Some utility fuctions used by the module are collected in this appendix. Their macro names can be seen in the below script.

⟨ Coding utilities 11a ⟩ ≡

```

""" Coding utilities """
⟨Generation of a random 3D point 12b⟩
⟨Generation of random 3D triangles 11b⟩
⟨Generation of random 3D quadrilaterals 12a⟩
⟨Generation of a single random triangle 12c⟩
⟨Containment boxes 13a⟩
⟨Transformation of a 3D box into an hexahedron 13b⟩
⟨Computation of the 1D centroid of a list of 3D boxes 14⟩
◇

```

Macro referenced in 6b.

Generation of random triangles The function `randomTriangles` returns the array `randomTriaArray` with a given number of triangles generated within the unit 3D interval. The `scaling` parameter is used to scale every such triangle, generated by three random points, that could be possibly located to far from each other, even at the distance of the diagonal of the unit cube.

The arrays `xs`, `ys` and `zs`, that contain the x, y, z coordinates of triangle points, are used to compute the minimal translation v needed to transport the entire set of data within the positive octant of the 3D space.

```

⟨Generation of random 3D triangles 11b⟩ ≡
""" Generation of random triangles """
def randomTriangles(numberOfTriangles=400,scaling=0.3):
    randomTriaArray = [rtriangle(scaling) for k in range(numberOfTriangles)]
    [xs,ys,zs] = TRANS(CAT(randomTriaArray))
    xmin, ymin, zmin = min(xs), min(ys), min(zs)
    v = array([-xmin,-ymin, -zmin])
    randomTriaArray = [[list(v1+v), list(v2+v), list(v3+v)] for v1,v2,v3 in randomTriaArray]
    return randomTriaArray
◇

```

Macro referenced in 11a.

Generation of random 3D quadrilaterals

```

⟨Generation of random 3D quadrilaterals 12a⟩ ≡
""" Generation of random 3D quadrilaterals """
def randomQuads(numberOfQuads=400,scaling=0.3):
    randomTriaArray = [rtriangle(scaling) for k in range(numberOfQuads)]
    [xs,ys,zs] = TRANS(CAT(randomTriaArray))
    xmin, ymin, zmin = min(xs), min(ys), min(zs)
    v = array([-xmin,-ymin, -zmin])
    randomQuadArray = [AA(list)([ v1+v, v2+v, v3+v, v+v2-v1+v3 ]) for v1,v2,v3 in randomTriaArray]
    return randomQuadArray
◇

```

Macro referenced in 11a.

Generation of a random 3D point A single random point, codified in floating point format, and with a fixed (quite small) number of digits, is returned by the `rpoint()` function, with no input parameters.

```

⟨ Generation of a random 3D point 12b ⟩ ≡
    """ Generation of a random 3D point """
    def rpoint():
        return eval( vcode([ random.random(), random.random(), random.random() ]) )
    ◇

```

Macro referenced in [11a](#).

Generation of a single random triangle A single random triangle, scaled about its centroid by the `scaling` parameter, is returned by the `rtriangle()` function, as a tuple of two random points in the unit square.

```

⟨ Generation of a single random triangle 12c ⟩ ≡
    """ Generation of a single random triangle """
    def rtriangle(scaling):
        v1,v2,v3 = array(rpoint()), array(rpoint()), array(rpoint())
        c = (v1+v2+v3)/3
        pos = rpoint()
        v1 = (v1-c)*scaling + pos
        v2 = (v2-c)*scaling + pos
        v3 = (v3-c)*scaling + pos
        return tuple(eval(vcode(v1))), tuple(eval(vcode(v2))), tuple(eval(vcode(v3)))
    ◇

```

Macro referenced in [11a](#).

Containment boxes Given as input a list `randomTriaArray` of pairs of 2D points, the function `containmentBoxes` returns, in the same order, the list of *containment boxes* of the input lines. A *containment box* of a geometric object of dimension d is defined as the minimal d -cuboid, equioriented with the reference frame, that contains the object. For a 2D line it is given by the tuple $(x1,y1,x2,y2)$, where $(x1,y1)$ is the point of minimal coordinates, and $(x2,y2)$ is the point of maximal coordinates.

```

⟨ Containment boxes 13a ⟩ ≡
    """ Containment boxes """
    def containmentBoxes(randomPointArray,qualifier=0):
        if len(randomPointArray[0])==2:
            boxes = [eval(vcode([min(x1,x2), min(y1,y2), min(z1,z2),
                                max(x1,x2), max(y1,y2), max(z1,z2)])))+[qualifier]
                    for ((x1,y1,z1),(x2,y2,z2)) in randomPointArray]
        elif len(randomPointArray[0])==3:
            boxes = [eval(vcode([min(x1,x2,x3), min(y1,y2,y3), min(z1,z2,z3),

```

```

        max(x1,x2,x3), max(y1,y2,y3), max(z1,z2,z3)))+[qualifier]
        for ((x1,y1,z1),(x2,y2,z2),(x3,y3,z3)) in randomPointArray]
    elif len(randomPointArray[0])==4:
        boxes = [eval(vcode([min(x1,x2,x3,x4), min(y1,y2,y3,y4), min(z1,z2,z3,z4),
            max(x1,x2,x3,x4), max(y1,y2,y3,y4), max(z1,z2,z3,z4)))+[qualifier]
            for ((x1,y1,z1),(x2,y2,z2),(x3,y3,z3),(x4,y4,z4)) in randomPointArray]
    return boxes

```

◇

Macro referenced in 11a.

Transformation of a 3D box into an hexahedron The transformation of a 2D box into a closed rectangular polyline, given as an ordered sequwncw of 2D points, is produced by the function `box2exa`

```

⟨ Transformation of a 3D box into an hexahedron 13b ⟩ ≡
    """ Transformation of a 3D box into an hexahedron """
    def box2exa(box):
        x1,y1,z1,x2,y2,z2,type = box
        verts = [[x1,y1,z1], [x1,y1,z2], [x1,y2,z1], [x1,y2,z2], [x2,y1,z1], [x2,y1,z2], [x2,y2,z1], [x2,y2,z2]]
        cell = [range(1,len(verts)+1)]
        return [verts,cell,None],type

    def lar2boxes(model,qualifier=0):
        V,CV = model
        boxes = []
        for k,cell in enumerate(CV):
            verts = [V[v] for v in cell]
            x1,y1,z1 = [min(coord) for coord in TRANS(verts)]
            x2,y2,z2 = [max(coord) for coord in TRANS(verts)]
            boxes += [eval(vcode([min(x1,x2),min(y1,y2),min(z1,z2),max(x1,x2),max(y1,y2),max(z1,z2)])+[qualifier]
            return boxes

```

◇

Macro referenced in 11a.

Computation of the 1D centroid of a list of 3D boxes The 1D centroid of a list of 3D boxes is computed by the function given below. The direction of computation (either x , y or z) is chosen depending on the value of the `coord` parameter.

```

⟨ Computation of the 1D centroid of a list of 3D boxes 14 ⟩ ≡
    """ Computation of the 1D centroid of a list of 3D boxes """
    def centroid(boxes,coord):
        delta,n = 0,len(boxes)
        ncoords = len(boxes[0])/2
        a = coord%ncoords
        b = a+ncoords

```

```
for box in boxes:
    delta += (box[a] + box[b])/2
return delta/n
```

◇

Macro referenced in [11a](#).

References

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