The drawing objects Technical details and regression checks

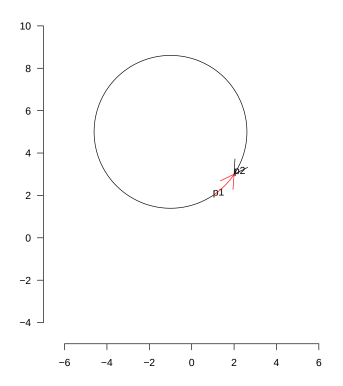
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17th July, 2009

1 The VDedgeSector object

A sector is a segment of a circle, defined by two points, together with the convention that a right-handed sector goes clockwise.

```
> nodeList <- list(p1 = matrix(1:2, ncol = 2), p2 = matrix(2:3,
      ncol = 2))
> centre = c(-1, 5)
> fromTheta <- .point.xy.to.theta(nodeList[["p1"]], centre)</pre>
> toTheta <- .point.xy.to.theta(nodeList[["p2"]], centre)</pre>
> 1h \leftarrow newEdgeSector(centre = c(-1, 5), hand = 1, from = "p1",
      to = "p2", fromTheta = fromTheta, toTheta = toTheta,
      radius = sqrt(13)
> lh <- .normalise.sector(lh)</pre>
> VD1 <- new("TissueDrawing", nodeList = nodeList)
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-7, 7), c(-5, 10))
> grid.xaxis()
> grid.yaxis()
> PlotNodes(VD1)
> xy <- .edge.to.xy(lh)</pre>
> grid.lines(xy[, 1], xy[, 2], default.units = "native",
      arrow = arrow())
> lh@hand <- -1
> xy <- .edge.to.xy(lh)</pre>
> grid.lines(xy[, 1], xy[, 2], default.units = "native",
      arrow = arrow(), gp = gpar(col = "red"))
```



We can also split VDedgeSectors

2 The TissueDrawing object

First we test constucting them from scratch.

```
> VD.nodeList <- list(p1 = matrix(1:2, ncol = 2), p2 = matrix(2:3,
      ncol = 2), p3 = matrix(c(-1, 0), ncol = 2))
> sectorfromto <- function(sector, from, to, nodeList) {
      sector@from <- from
      sector@to <- to
      from.point <- nodeList[[from]]</pre>
      sector@fromTheta <- .point.xy.to.theta(from.point,</pre>
          sector@centre)
      sector@toTheta <- .point.xy.to.theta(nodeList[[to]],</pre>
+
+
          sector@centre)
      sector <- .normalise.sector(sector)</pre>
+ }
> centre = c(-1, 5)
> fromTheta <- .point.xy.to.theta(nodeList[["p1"]], centre)</pre>
> toTheta <- .point.xy.to.theta(nodeList[["p2"]], centre)</pre>
> 1h < -newEdgeSector(centre = c(-1, 5), hand = 1, fromTheta = fromTheta,
      toTheta = toTheta, radius = sqrt(13))
> lh <- sectorfromto(lh, "p1", "p2", VD.nodeList)
> centre = c(4, 0)
> fromTheta <- .point.xy.to.theta(nodeList[["p1"]], centre)</pre>
> toTheta <- .point.xy.to.theta(nodeList[["p2"]], centre)</pre>
> rh <- newEdgeSector(centre = c(4, 0), hand = 1, fromTheta = fromTheta,
      toTheta = toTheta, radius = sqrt(13))
> el <- newEdgeLines(from = "p1", to = "p3", xy = matrix(c(1,
      2, -0.5, 0, -1, 0), ncol = 2, byrow = T)
> VD.edgeList <- list(`p1|p2|1` = sectorfromto(lh, "p1",
      "p2", VD.nodeList), `p2|p1|1` = sectorfromto(lh,
      "p2", "p1", VD.nodeList), `p1|p2|2` = sectorfromto(rh,
      "p1", "p2", VD.nodeList), `p2|p1|2` = sectorfromto(rh, "p2", "p1", VD.nodeList), `p1|p3|3` = e1, `p3|p1|3` = newEdgeLines(from = "p3",
      to = "p1", xy = matrix(c(-1, 0, 1, 2), ncol = 2,
          byrow = T)))
> VD.faceList <- list(`100` = c("p1|p2|1", "-p1|p2|2"),
      `110` = c("p1|p2|2", "p2|p1|1"), `010` = c("p2|p1|2",
           "-p2|p1|1"), `001` = c("p1|p3|3", "p3|p1|3"),
      \label{eq:defDarkMatter} \textit{DarkMatter} = c("-p3|p1|3", "-p1|p3|3", "-p2|p1|2",
          "-p1|p2|1"))
> VD.setList <- list(`1` = c("p1|p2|1", "p2|p1|1"), `2` = c("p1|p2|2",
       "p2|p1|2"), `3` = c("p1|p3|3", "p3|p1|3"))
> VD.faceSignature <- lapply(names(VD.faceList), function(x) {
+
+ })
> names(VD.faceSignature) <- names(VD.faceList)</pre>
> VD <- new("TissueDrawing", nodeList = VD.nodeList, edgeList = VD.edgeList,
      setList = VD.setList, faceList = VD.faceList, faceSignature = VD.faceSignature)
> .validateDrawing(VD)
```

```
Validating a drawing on 3 sets.....done
> VD
> .checkPointOnEdge(edge = VD@edgeList[["p1|p2|1"]], point.xy = VD@nodeList[["p1"]])
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-7, 7), c(-5, 10))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(VD)
> PlotSetBoundaries(VD)
```

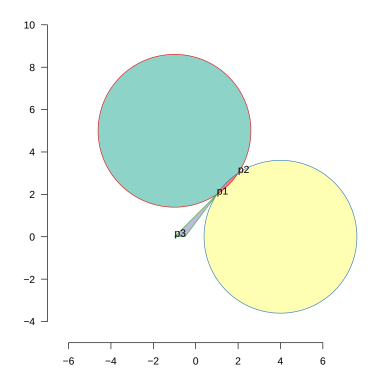
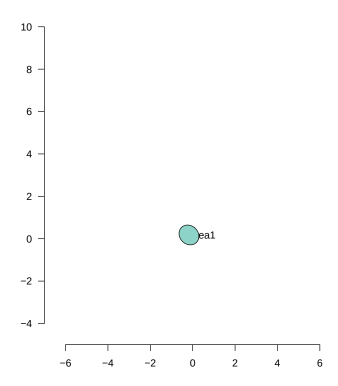


Figure 1: Constructing TissueDrawing objects from scratch

2.1 Ellipses

> PlotNodes(VD)

Ellipses could be coped with specially by finding roots of quartics, but don't bother and just generate them as polygons



```
phi, e, a, Set = 4, dx = dx)
> TM <- VE[[1]]
 TM2 <- addSetToDrawing(TM, VE[[2]], set2Name = paste("Set",</pre>
      2, sep = ""))
> TM3 <- addSetToDrawing(TM2, VE[[3]], set2Name = paste("Set",
      3, sep = "")
> TM4 <- addSetToDrawing(TM3, VE[[4]], set2Name = paste("Set",
      4, sep = "")
> .validateDrawing(TM4)
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-10, 10), c(-8, 10))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM4)
> PlotSetBoundaries(TM4, gp = gpar(1wd = 2, col = c("red",
      "blue", "green", "yellow")))
> .PlotFaceNames.TissueDrawing(TM4)
```

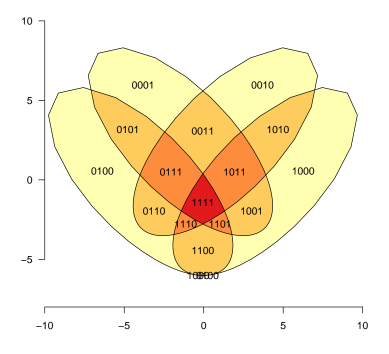


Figure 2: Constructing TissueDrawing objects from scratch

3 Injecting points and edges

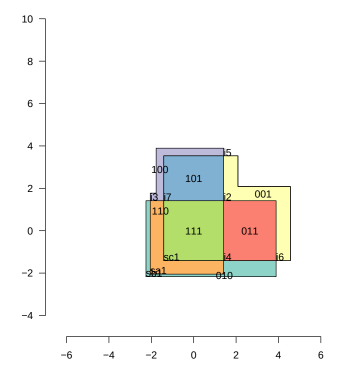
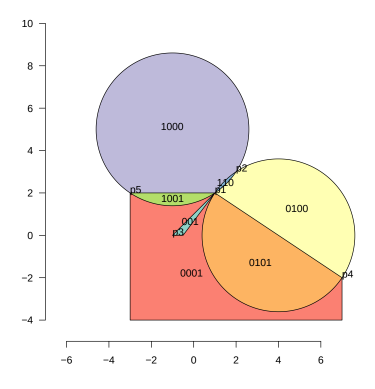


Figure 3: Injecting points

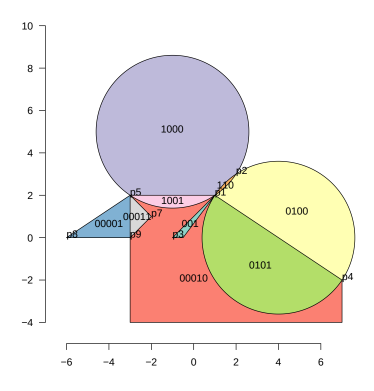
Then we try injecting single edges

```
> VD6 <- injectEdge(drawing = VD6, newEdgeList = list(`p5|p1|4` = p5p1.line),
      set2Name = "4", addToList = FALSE)
> VD6 <- injectEdge(drawing = VD6, newEdgeList = list(`p4|p5|4` = p4p5.line),
      set2Name = "4", addToList = FALSE)
 .is.face.within.set(drawing = VD6, faceName = "0101",
      setName = "2")
> .is.face.within.set(drawing = VD6, faceName = "1000",
      setName = "2")
 .is.face.within.set(drawing = VD6, faceName = "0001",
      setName = "2")
> VD6
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-7, 7), c(-5, 10))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(VD6)
> PlotSetBoundaries(VD6, gp = gpar(lwd = 2, col = c("red",
      "blue", "green", "black")))
> .PlotFaceNames.TissueDrawing(VD6)
> PlotNodes(VD6)
```



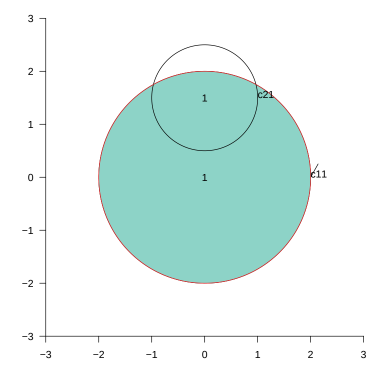
And now injecting edges of multiple points

```
> VD8 <- VD6
> p7 \leftarrow matrix(c(-2, 1), ncol = 2)
> rownames(p7) <- "p7"
> VD8@nodeList[["p7"]] <- p7</pre>
> p8 \leftarrow matrix(c(-6, 0), ncol = 2)
> rownames(p8) <- "p8"
> VD8@nodeList[["p8"]] <- p8</pre>
> p9 \leftarrow matrix(c(-3, 0), ncol = 2)
> rownames(p9) <- "p9"
> VD8@nodeList[["p9"]] <- p9</pre>
> p5p7.line <- newEdgeLines(from = "p5", to = "p7", xy = matrix(c(-3, -3))
      2, -2, 1), ncol = 2, byrow = T))
> p7p9.line \leftarrow newEdgeLines(from = "p7", to = "p9", xy = matrix(c(-2, p7p9))
      1, -3, 0), ncol = 2, byrow = T)
> p9p8.line \leftarrow newEdgeLines(from = "p9", to = "p8", xy = matrix(c(-3, p9p8.line)))
      0, -6, 0), ncol = 2, byrow = T)
> p8p5.line <- newEdgeLines(from = "p8", to = "p5", xy = matrix(c(-6,
      0, -3, 2), ncol = 2, byrow = T))
> VD8@edgeList[["p5|p7|5"]] <- p5p7.line
> VD8@edgeList[["p7|p9|5"]] <- p7p9.line</pre>
> VD8@edgeList[["p9|p8|5"]] <- p9p8.line
> VD8@edgeList[["p8|p5|5"]] <- p8p5.line
> VD8@setList[["5"]] <- c("p5|p7|5", "p7|p9|5", "p9|p8|5",
      "p8|p5|5")
> VD8@edgeList[["p4|p5|4"]]@xy
> VD8 <- injectPoint(drawing = VD8, edgeName = "p4|p5|4",
      newPoint = VD8@nodeList[["p9"]])
> VD8@edgeList[["p9|p5|4"]]@xy
> VD8@edgeList[["p4|p9|4"]]@xy
> VD8 <- injectEdge(drawing = VD8, newEdgeList = VD8@edgeList[c("p5|p7|5",
      "p7|p9|5")], set2Name = "5", addToList = FALSE)
> VD8 <- injectEdge(drawing = VD8, newEdgeList = VD8@edgeList[c("p9|p8|5",
      "p8|p5|5")], set2Name = "5", addToList = FALSE)
```



4 Making a simple drawing from a circle

```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> xy <- .edge.to.xy(VDC1@edgeList[[1]])
> grid.lines(xy[, 1], xy[, 2], default.units = "native",
+ arrow = arrow())
> PlotFaces(VDC1)
> PlotFaces(VDC2, gp = gpar(fill = "red"))
> PlotSetBoundaries(VDC1)
> .PlotFaceNames.TissueDrawing(VDC1)
> PlotNodes(VDC2)
> .PlotFaceNames.TissueDrawing(VDC2)
```



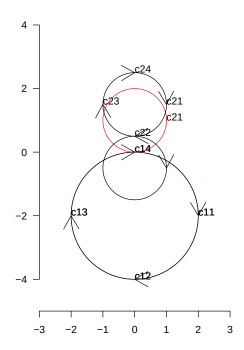
5 Circles

```
> r = 0.6
> d = 0.4
> angles <- pi/2 - c(0, 2 * pi/3, 4 * pi/3)
> x <- d * cos(angles)
> y <- d * sin(angles)
> r <- rep(r, 3)</pre>
```

6 Non overlapping circles

```
> centre.xy <- c(0, -2)
> VDC1 <- newTissueFromCircle(centre.xy, radius = 2, Set = 1,
+ nodes = 4)
> VDC2 <- newTissueFromCircle(centre.xy + c(0, 3.5), radius = 1,
+ Set = 2, nodes = 4)
> TN2 <- addSetToDrawing(VDC1, VDC2)
> VDC3 <- newTissueFromCircle(c(0, -0.5), radius = 1, Set = 3)
> .validateDrawing(TN2)
> centre.xy <- c(0, -2)
> VDC1b <- newTissueFromCircle(centre.xy, radius = 2, Set = 1,
+ nodes = 4)
> VDC2b <- newTissueFromCircle(centre.xy + c(0, 3), radius = 1,
+ Set = 2)
> TN2b <- (addSetToDrawing(VDC1b, VDC2b))
> try(.validateDrawing(TN2b))
```

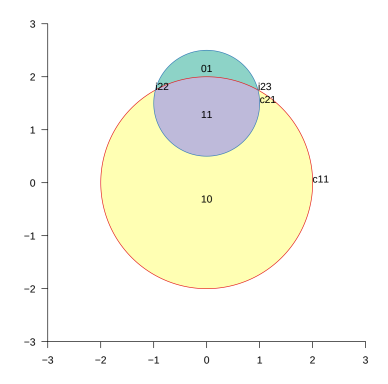
```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-5, 5))
> grid.xaxis()
> grid.yaxis()
> shoar <- function(TN2) {</pre>
      for (ix in 1:length(TN2@edgeList)) {
          xy <- .edge.to.xy(TN2@edgeList[[ix]])</pre>
          grid.lines(xy[, 1], xy[, 2], default.units = "native",
              arrow = arrow())
+ }
> PlotSetBoundaries(VDC1b)
> PlotNodes(VDC1b)
> PlotSetBoundaries(VDC2b)
> PlotNodes(VDC2b)
> shoar(VDC1)
> shoar(VDC2)
> shoar(VDC3)
> PlotNodes(TN2)
```



7 Check for the intersection of two edges

```
> centre.xy <- c(0, 0)
> VDC1 <- newTissueFromCircle(centre.xy, radius = 2, Set = 1)
> renameFaces(VDC1, oldName = .faceNames(VDC1, onlyVisible = TRUE),
      "1")
> VDC2 <- newTissueFromCircle(centre.xy + c(0, 1.5), radius = 1,
      Set = 2
> edge1 <- VDC1@edgeList[[1]]</pre>
> edge2 <- VDC2@edgeList[[1]]</pre>
> .findIntersection(edge1, edge2)
> edge1 <- VD8@edgeList[["p1|p4|4"]]</pre>
> edge2 <- VDC2@edgeList[[1]]</pre>
> .findIntersection(edge1, edge2)
> edge1 <- VD8@edgeList[["p1|p4|4"]]</pre>
> edge2 <- VD8@edgeList[["p2|p4|2"]]</pre>
> .findIntersection(edge1, edge2)
> .find.point.within.face(drawing = VD8, faceName = "1001")
> .is.point.within.face(VD8, "DarkMatter", p7)
> .is.point.within.face(VD8, "DarkMatter", matrix(c(-100,
      100), ncol = 2))
> edge1 <- VD8@edgeList[["p1|p4|4"]]</pre>
> edge2 <- VD8@edgeList[["p1|p3|3"]]</pre>
> .findIntersection(edge1, edge2)
> drawing1 <- VDC1
> drawing2 <- VDC2
> VM <- addSetToDrawing(drawing1 = VDC1, drawing2 = VDC2,
      set2Name = "Set2")
> .validateDrawing(VM)
```

```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(VM)
> PlotSetBoundaries(VM)
> .PlotFaceNames.TissueDrawing(VM)
> PlotNodes(VM)
```



8 addSetToDrawing two polygons

```
> d <- 1

> s1 <- 0.7

> s2 <- 0.6

> d <- 0.9146274

> s1 <- 2.44949

> s2 <- 2.645751

> 11 <- -d/2 - s1/2

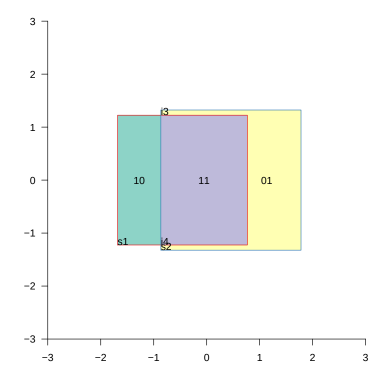
> 12 <- d/2 - s2/2

> r1 <- -d/2 + s1/2

> r2 <- d/2 + s2/2

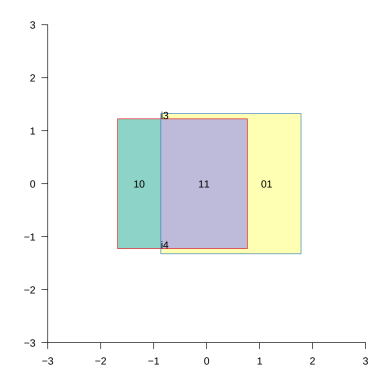
> poly.1 <- matrix(c(11, -s1/2, 11, s1/2, r1, s1/2, r1, + -s1/2), ncol = 2, byrow = TRUE)
```

```
> rownames(poly.1) <- paste("s", 1:4, sep = "")
> poly.2 <- matrix(c(12, -s2/2, 12, s2/2, r2, s2/2, r2,
      -s2/2), ncol = 2, byrow = TRUE)
> rownames(poly.2) <- paste("s", 2:5, sep = "")
> VDP1 <- newTissueFromPolygon(points.xy = poly.1, Set = 1)
> VDP2 <- newTissueFromPolygon(points.xy = poly.2, Set = 2)
> TM <- addSetToDrawing(drawing1 = VDP1, drawing2 = VDP2,
      set2Name = "Set2")
> .validateDrawing(TM)
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM)
> PlotSetBoundaries(TM)
> .PlotFaceNames.TissueDrawing(TM)
> PlotNodes(TM)
```



> TMR <- remove.nonintersectionpoints(drawing = TM)
> .validateDrawing(TMR)

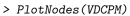
```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TMR)
> PlotSetBoundaries(TMR)
> .PlotFaceNames.TissueDrawing(TMR)
> PlotNodes(TMR)
```

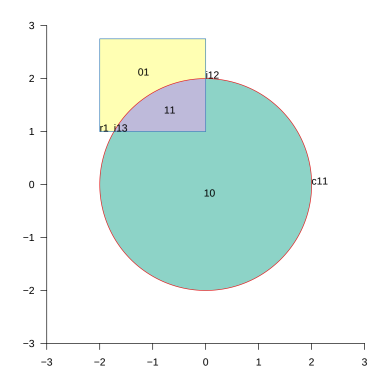


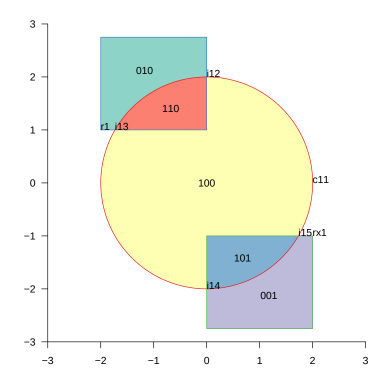
9 addSetToDrawing a polygon and a circle

> .validateDrawing(VDCPM)

```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(VDCPM)
> PlotSetBoundaries(VDCPM)
> .PlotFaceNames.TissueDrawing(VDCPM)
```

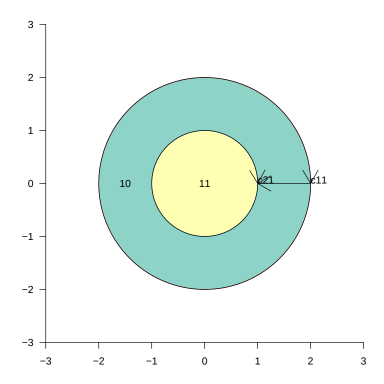






10 Invisible edges

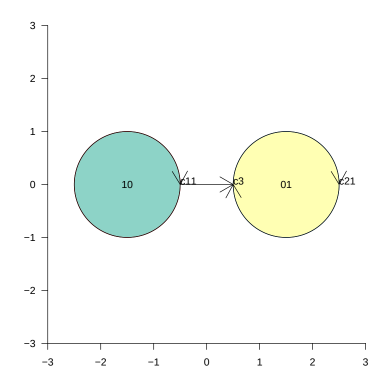
```
> centre.xy <- c(0, 0)
> VDC3 <- newTissueFromCircle(centre.xy, radius = 2, Set = 1)
> VDC4 <- newTissueFromCircle(centre.xy, radius = 1, Set = 2)
> VDI <- addSetToDrawing(drawing1 = VDC3, drawing2 = VDC4,
+ set2Name = "Set2")
> .validateDrawing(VDI)
```



The code only attemtps to inject invisible edges between known points, so we have to give the algorithm a hint by inserting such known points in the right place

```
> centre.xy <- c(-1.5, 0)
> VDC5 <- newTissueFromCircle(centre.xy, radius = 1, Set = 1)
> VDC6 <- newTissueFromCircle(centre.xy + c(3, 0), radius = 1,
+ Set = 2)
> VDC6 <- injectPoint(VDC6, "c21/c21/2", newPoint = matrix(c(0.5,
+ 0), ncol = 2, dimnames = list("c3")))
> VDO <- addSetToDrawing(drawing1 = VDC5, drawing2 = VDC6,
+ set2Name = "Set2")</pre>
```

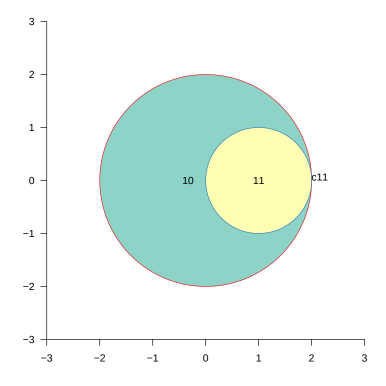
```
> .validateDrawing(VDO)
```



11 Tangents

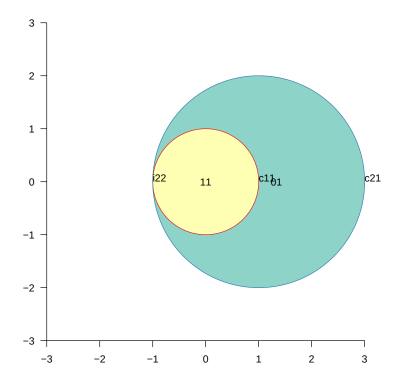
```
+ set2Name = "Set2")
> .validateDrawing(VDT)

> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(VDT)
> PlotSetBoundaries(VDT)
> .PlotFaceNames.TissueDrawing(VDT)
> PlotNodes(VDT)
```

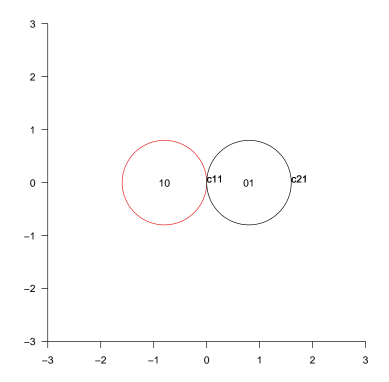


```
> centre.xy <- c(0, 0)
> VDC9 <- newTissueFromCircle(centre.xy, radius = 1, Set = 1)
> VDC10 <- newTissueFromCircle(centre.xy + c(1, 0), radius = 2,
+ Set = 2)
> VDT2 <- addSetToDrawing(drawing1 = VDC9, drawing2 = VDC10,
+ set2Name = "Set2")
> .validateDrawing(VDT2)
```

```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(VDT2)
> PlotSetBoundaries(VDT2)
> .PlotFaceNames.TissueDrawing(VDT2)
> PlotNodes(VDT2)
```



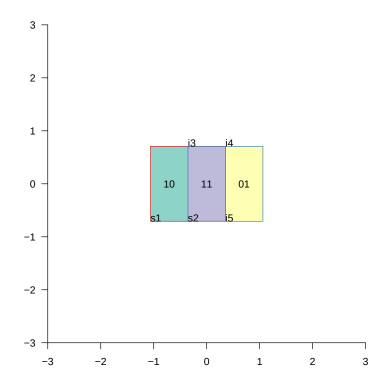
```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotSetBoundaries(VDC1)
> PlotSetBoundaries(VDC2, gp = gpar(col = "red"))
> PlotNodes(VDC1)
> PlotNodes(VDC2)
> .PlotFaceNames.TissueDrawing(VDT)
> PlotNodes(VDT)
```



```
> 11 <- -1.06066
> r1 <- 0.3535534
> 12 <- -0.3535534
> r2 <- 1.06066
> s1 <- 1.414214
> s2 <- 1.414214
> poly.1 <- matrix(c(11, -s1/2, 11, s1/2, r1, s1/2, r1, + -s1/2), ncol = 2, byrow = TRUE)
> rownames(poly.1) <- paste("s", 1:4, sep = "")
> poly.2 <- matrix(c(12, -s2/2, 12, s2/2, r2, s2/2, r2, + -s2/2), ncol = 2, byrow = TRUE)
> rownames(poly.2) <- paste("s", 2:5, sep = "")
> VDP1 <- newTissueFromPolygon(points.xy = poly.1, Set = 1)</pre>
```

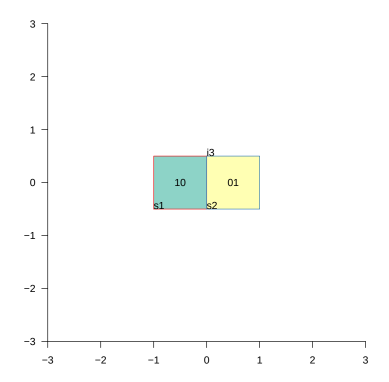
```
> VDP2 <- newTissueFromPolygon(points.xy = poly.2, Set = 2)
> TM <- addSetToDrawing(drawing1 = VDP1, drawing2 = VDP2,
+ set2Name = "Set2")

> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM)
> PlotSetBoundaries(TM)
> .PlotFaceNames.TissueDrawing(TM)
> PlotNodes(TM)
```



```
+ -s2/2), ncol = 2, byrow = TRUE)
> rownames(poly.2) <- paste("s", 2:5, sep = "")
> VDP3 <- newTissueFromPolygon(points.xy = poly.1, Set = 1)
> VDP4 <- newTissueFromPolygon(points.xy = poly.2, Set = 2)
> TM3 <- addSetToDrawing(drawing1 = VDP3, drawing2 = VDP4,
+ set2Name = "Set2")

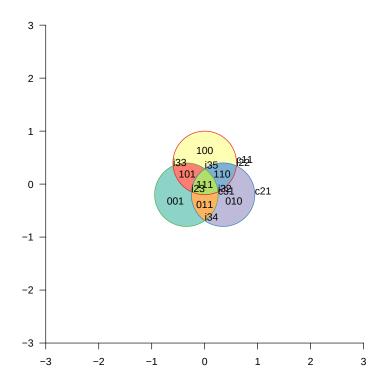
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM3)
> PlotSetBoundaries(TM3)
> PlotFaceNames.TissueDrawing(TM3)
> PlotNodes(TM3)
```



12 Three circles

```
> r <- 0.6
> d <- 0.4
> angles <- pi/2 - c(0, 2 * pi/3, 4 * pi/3)
> x <- d * cos(angles)
> y <- d * sin(angles)</pre>
```

```
> r \leftarrow rep(r, 3)
> centres <- matrix(c(x, y), ncol = 2, byrow = FALSE)
> VDC1 <- newTissueFromCircle(centres[1, ], radius = r[1],
      Set = 1)
> VDC2 <- newTissueFromCircle(centres[2, ], radius = r[2],
      Set = 2)
> TM3 <- addSetToDrawing(drawing1 = VDC1, drawing2 = VDC2,
      set2Name = "Set2")
 VDC3 <- newTissueFromCircle(centres[3, ], radius = r[3],
      Set = 3)
> TM3 <- addSetToDrawing(drawing1 = TM3, drawing2 = VDC3,
      set2Name = "Set3")
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM3)
> PlotSetBoundaries(TM3)
> .PlotFaceNames.TissueDrawing(TM3)
> PlotNodes(TM3)
```



13 Three circles

```
> r \leftarrow c(1.261566, 0.977205, 1.492705)

> x \leftarrow c(0, 1.350138, -1.086542)

> y \leftarrow c(1.2615663, -0.8066661, -0.4028718)

> centres \leftarrow matrix(c(x, y), ncol = 2, byrow = FALSE)

> VDC1 \leftarrow newTissueFromCircle(centres[1, ], radius = r[1],

+ Set = 1, nodes = 5)

> VDC2 \leftarrow newTissueFromCircle(centres[2, ], radius = r[2],

+ Set = 2, nodes = 5)

> TM \leftarrow addSetToDrawing(drawing1 = VDC1, drawing2 = VDC2,

+ set2Name = "Set2")

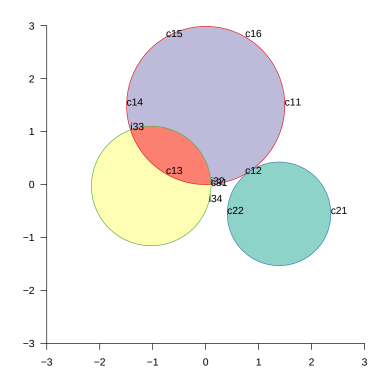
> VDC3 \leftarrow newTissueFromCircle(centres[3, ], radius = r[3],

+ Set = 3)

> TM \leftarrow addSetToDrawing(drawing1 = TM, drawing2 = VDC3,

+ set2Name = "Set3")
```

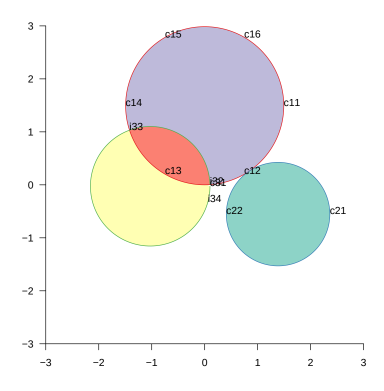
```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM3)
> PlotSetBoundaries(VDC1)
> PlotSetBoundaries(VDC2)
> PlotNodes(VDC1)
> PlotNodes(VDC2)
> .PlotFaceNames.TissueDrawing(TM3)
> PlotNodes(TM3)
```



14 Three circles

```
+ set2Name = "Set2")
> VDC3 <- newTissueFromCircle(centres[3, ], radius = r[3],
+ Set = 3)
> TM3 <- addSetToDrawing(drawing1 = TM, drawing2 = VDC3,
+ set2Name = "Set3")
> TV3 <- .merge.faces.invisibly.split(TM3)

> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-3, 3), c(-3, 3))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TV3)
> PlotSetBoundaries(TV3)
> PlotNodes(TM3)
```



15 Triangles

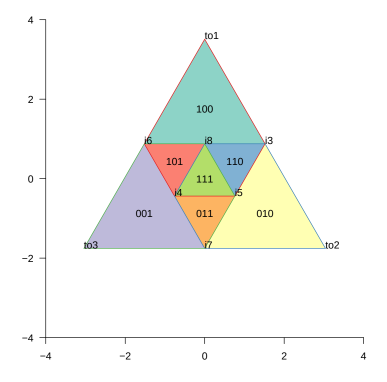
```
> .inscribetriangle.feasible <- function(wghts) {
+     w0 <- 1 - sum(wghts)
+     stopifnot(all(wghts <= 1) & all(wghts >= 0) & w0 >=
+         0)
+     wa <- wghts[1]
+     wb <- wghts[2]</pre>
```

```
wc <- wghts[3]</pre>
      Delta \leftarrow w0^2 - 4 * wa * wb * wc
+
+
      return(Delta >= 0)
+ }
> .inscribetriangle.compute <- function(wghts) {</pre>
      wa <- wghts[1]</pre>
+
+
      wb <- wghts[2]
      wc <- wghts[3]
      stopifnot(.inscribetriangle.feasible(wghts))
      pa <- (1 - wc)
      pb < - (wb + wc - wa - 1)
      pc <- wa * (1 - wb)
      sc <- if (wa > 0) {
           (-pb - sqrt(pb^2 - 4 * pa * pc))/(2 * pa)
+
+
      else if (wb + wc < 1) {
          (1 - wb - wc)/(1 - wc)
+
+
      else {
+
           0
      }
+
      sb <- if (sc > 0) {
           1 - wa/sc
      }
+
      else {
           wc/(1 - wb)
+
      sa \leftarrow wb/(1 - sc)
+
      c(sc, sa, sb)
+ }
> .inscribetriangle.inscribe <- function(xy, wghts) {</pre>
+
      scalef <- NA
+
      isfeasible <- .inscribetriangle.feasible(wghts)</pre>
      if (!isfeasible) {
           scalef <- 4 * wghts[1] * wghts[2] * wghts[3]/(1 -</pre>
+
               sum(wghts))^2
+
           scalef <- scalef^(1/3)</pre>
           wghts <- wghts/(scalef * 1.001)</pre>
           isfeasible <- .inscribetriangle.feasible(wghts)</pre>
+
           stopifnot(!isfeasible)
+
      }
      if (!isfeasible)
+
           return(list(feasible = FALSE))
      scab <- .inscribetriangle.compute(wghts)</pre>
+
      inner.xy <- (1 - scab) * xy + scab * (xy[c(2, 3, 3)])
      return(list(feasible = TRUE, inner.xy = inner.xy,
           scalef = scalef))
+ }
```

```
> WeightUniverse <- 18
> WeightVisible <- 16
> WeightInvisible <- WeightUniverse - WeightVisible
> w0ratio <- WeightInvisible/WeightVisible</pre>
> wa <- 0.25
> wb <- 0.25
> wc <- 0.25
> outer.weights <- c(wa, wb, wc)
> outer.innerw <- 1 - sum(outer.weights)</pre>
> outer.inner.ratios <- outer.weights/outer.innerw
> outer.feasible <- .inscribetriangle.feasible(outer.weights)
> wab <- 0.0625
> wbc <- 0.0625
> wca <- 0.0625
> wabc <- 0.0625
> inner.weights <- c(wab, wbc, wca)
> inner.innerw <- wabc
> sf <- (sum(inner.weights) + inner.innerw)</pre>
> Weight.Inner <- sf * WeightVisible
> if (sf > 0) {
      inner.weights <- inner.weights/sf</pre>
      inner.feasible <- .inscribetriangle.feasible(inner.weights)</pre>
+ } else {
      inner.feasible <- FALSE
+ }
> side <- sqrt(4 * WeightVisible/(3 * sqrt(3)))</pre>
> angles <- pi/2 - c(0, 2 * pi/3, 4 * pi/3)
> outer.xy <- t(sapply(angles, function(a) c(x = side *
      cos(a), y = side * sin(a))))
> inner <- .inscribetriangle.inscribe(outer.xy, wghts = outer.weights)</pre>
> inner.xy <- inner$inner.xy</pre>
> innest <- .inscribetriangle.inscribe(inner.xy, wghts = inner.weights)</pre>
> innest.xy = innest$inner.xy
> outest.xy <- outer.xy * sqrt(1 + w0ratio)</pre>
> rownames(outer.xy) <- paste("to", 1:3, sep = "")</pre>
> rownames(inner.xy) <- paste("ti", 1:3, sep = "")</pre>
> rownames(innest.xy) <- paste("tt", 1:3, sep = "")</pre>
> outline.a.xy <- do.call(rbind, list(outer.xy[1, , drop = FALSE],</pre>
      inner.xy[1, , drop = FALSE], innest.xy[1, , drop = FALSE],
      innest.xy[2, , drop = FALSE], inner.xy[3, , drop = FALSE]))
> outline.b.xy <- do.call(rbind, list(outer.xy[2, , drop = FALSE],
      inner.xy[2, , drop = FALSE], innest.xy[2, , drop = FALSE],
      innest.xy[3, , drop = FALSE], inner.xy[1, , drop = FALSE]))
> outline.c.xy <- do.call(rbind, list(outer.xy[3, , drop = FALSE],
      inner.xy[3, , drop = FALSE], innest.xy[3, , drop = FALSE],
      innest.xy[1, , drop = FALSE], inner.xy[2, , drop = FALSE]))
> VDP1 <- newTissueFromPolygon(points.xy = outline.a.xy,
      Set = 1)
> VDP2 <- newTissueFromPolygon(points.xy = outline.b.xy,
      Set = 2
```

```
> VDP3 <- newTissueFromPolygon(points.xy = outline.c.xy,
+     Set = 3)
> TMT <- addSetToDrawing(drawing1 = VDP1, drawing2 = VDP2,
+     set2Name = "Set2")
> TMT <- addSetToDrawing(drawing1 = TMT, drawing2 = VDP3,
+     set2Name = "Set3")

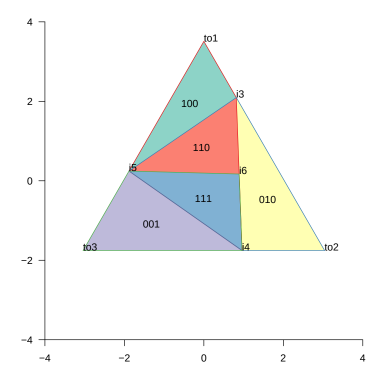
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-4, 4), c(-4, 4))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TMT)
> PlotSetBoundaries(TMT)
> PlotFaceNames.TissueDrawing(TMT)
> PlotNodes(TMT)
```



```
> WeightUniverse <- 18
> WeightVisible <- 16
> WeightInvisible <- WeightUniverse - WeightVisible
> w0ratio <- WeightInvisible/WeightVisible
> wa <- 0.1666666667
> wb <- 0.25
> wc <- 0.25
> outer.weights <- c(wa, wb, wc)</pre>
```

```
> outer.innerw <- 1 - sum(outer.weights)</pre>
> outer.inner.ratios <- outer.weights/outer.innerw
> outer.feasible <- .inscribetriangle.feasible(outer.weights)</pre>
> wab <- 0.16666667
> wbc <- 0
> wca <- 0
> wabc <- 0.16666667
> inner.weights <- c(wab, wbc, wca)
> inner.innerw <- wabc
> sf <- (sum(inner.weights) + inner.innerw)</pre>
> Weight.Inner <- sf * WeightVisible
> if (sf > 0) {
      inner.weights <- inner.weights/sf</pre>
      inner.feasible <- .inscribetriangle.feasible(inner.weights)</pre>
+ } else {
      inner.feasible <- FALSE
+ }
> side <- sqrt(4 * WeightVisible/(3 * sqrt(3)))</pre>
> angles <- pi/2 - c(0, 2 * pi/3, 4 * pi/3)
> outer.xy <- t(sapply(angles, function(a) c(x = side *
      cos(a), y = side * sin(a))))
> inner <- .inscribetriangle.inscribe(outer.xy, wghts = outer.weights)</pre>
> inner.xy <- inner$inner.xy</pre>
> innest <- .inscribetriangle.inscribe(inner.xy, wghts = inner.weights)</pre>
> innest.xy = innest$inner.xy
> outest.xy <- outer.xy * sqrt(1 + w0ratio)</pre>
> rownames(outer.xy) <- paste("to", 1:3, sep = "")</pre>
> rownames(inner.xy) <- paste("ti", 1:3, sep = "")</pre>
> rownames(innest.xy) <- paste("tt", 1:3, sep = "")</pre>
> outline.a.xy <- do.call(rbind, list(outer.xy[1, , drop = FALSE],
      inner.xy[1, , drop = FALSE], innest.xy[1, , drop = FALSE],
      innest.xy[2, , drop = FALSE], inner.xy[3, , drop = FALSE]))
> outline.b.xy <- do.call(rbind, list(outer.xy[2, , drop = FALSE],
      inner.xy[2, , drop = FALSE], innest.xy[2, , drop = FALSE],
      innest.xy[3, , drop = FALSE], inner.xy[1, , drop = FALSE]))
> outline.c.xy <- do.call(rbind, list(outer.xy[3, , drop = FALSE],
      inner.xy[3, , drop = FALSE], innest.xy[3, , drop = FALSE],
      innest.xy[1, , drop = FALSE], inner.xy[2, , drop = FALSE]))
> VDP1 <- newTissueFromPolygon(points.xy = outline.a.xy,
      Set = 1)
> VDP2 <- newTissueFromPolygon(points.xy = outline.b.xy,
> VDP3 <- newTissueFromPolygon(points.xy = outline.c.xy,
      Set = 3
> TMT <- addSetToDrawing(drawing1 = VDP1, drawing2 = VDP2,
     set2Name = "Set2")
> TMT <- addSetToDrawing(drawing1 = TMT, drawing2 = VDP3,
      set2Name = "Set3")
```

```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-4, 4), c(-4, 4))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TMT)
> PlotSetBoundaries(TMT)
> .PlotFaceNames.TissueDrawing(TMT)
> PlotNodes(TMT)
```



16 Three squares

```
> ss1 < -c(-2.04988805276466, 1.4142135623731, 1.41421356237309, +c.77228856812726, -1.77228856812726, -2.04988805276466, -2.04988805276466, 3.8936089116869, 3.8936089116869, 1.77228856812726, 1.77228856812726) > <math>ss2 < -c(-2.25237500351774, 3.88908729652601, 3.88908729652601, -2.25237500351774, -2.16799518941608, -2.16799518941608, +1.4142135623731, 1.41421356237309) > <math>ss3 < -c(-1.4142135623731, 4.56252232622749, 4.56252232622749, 2.08764859207457, 2.08764859207457, -1.4142135623731, -1.4142135623731, 2.08764859207457, +2.08764859207457, 3.53553390593274, 3.53553390593274) > SS1 < -matrix(ss1, ncol = 2, byrow = FALSE) > rownames(SS1) < -paste("sa", 1:6, sep = "")
```

```
> SS2 <- matrix(ss2, ncol = 2, byrow = FALSE)
> rownames(SS2) <- paste("sb", 1:4, sep = "")
> SS3 <- matrix(ss3, ncol = 2, byrow = FALSE)
> rownames(SS3) <- paste("sc", 1:6, sep = "")
> VDP1 <- newTissueFromPolygon(points.xy = SS1, Set = 1)
> VDP2 <- newTissueFromPolygon(points.xy = SS2, Set = 2)
> VDP3 <- newTissueFromPolygon(points.xy = SS3, Set = 3)
> TM <- addSetToDrawing(drawing1 = VDP1, drawing2 = VDP2,
      set2Name = "Set2")
> TM <- addSetToDrawing(drawing1 = TM, drawing2 = VDP3,
      set2Name = "Set3")
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-7, 7), c(-5, 10))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM)
> PlotSetBoundaries(TM, gp = gpar(lwd = 2, col = c("green",
      "red")))
> PlotNodes(TM)
> .PlotFaceNames.TissueDrawing(TM)
> PlotSetBoundaries(VDP3, gp = gpar(lwd = 2, col = c("green")))
```

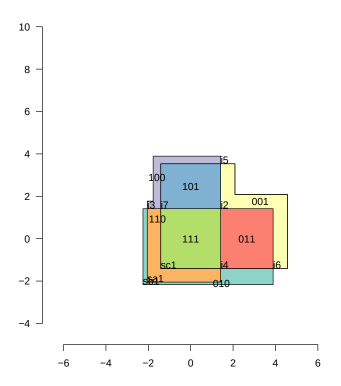


Figure 4: Injecting points

17 Noncontigous subsets

```
> px1 \leftarrow matrix(c(-5, -3, -5, 3, 5, 3, 5, -3), ncol = 2,
      byrow = TRUE)
> rownames(px1) <- paste("pa", 1:nrow(px1), sep = "")</pre>
 px2 \leftarrow matrix(c(-3, -5, -3, 5, 3, 5, 3, -5), ncol = 2,
      byrow = TRUE)
> rownames(px2) <- paste("pb", 1:nrow(px2), sep = "")</pre>
> VX1 <- newTissueFromPolygon(px1, Set = 1)
> VX2 <- newTissueFromPolygon(px2, Set = 2)
> TM <- addSetToDrawing(VX1, VX2, set2Name = "Set2")
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-10, 10), c(-10, 10))
> grid.xaxis()
> grid.yaxis()
> PlotNodes(TM)
> PlotSetBoundaries(TM, gp = gpar(lwd = 2, col = c("green",
      "red", "blue")))
> .PlotFaceNames.TissueDrawing(TM)
```

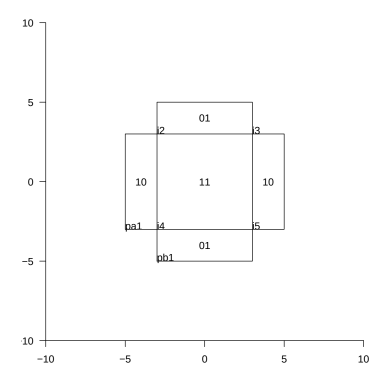


Figure 5: Injecting points

18 Ellipses

```
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-10, 10), c(-10, 10))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM)
> PlotSetBoundaries(E[[2]], gp = gpar(lwd = 2, col = c("red", "red", "blue")))
> PlotNodes(TM)
> .PlotFaceNames.TissueDrawing(TM)
> PlotSetBoundaries(TM, gp = gpar(lwd = 2, col = c("green")))
```

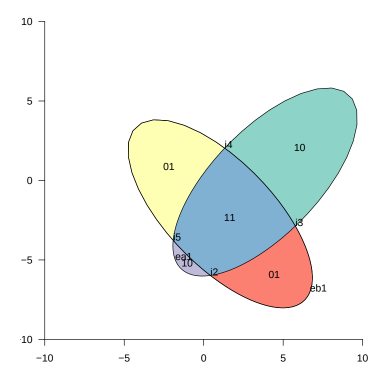


Figure 6: Injecting points

```
> phi < 0.8

> dex < 1.7

> dey < 2.5

> a < 7.6

> e < 0.9

> x0 < (-0.9, -5)

> dx < 0.1

> E(-1ist())

> E[[1]] < -newTissueFromEllipse(f1 = x0 + c(0, 0), dx = dx, + phi = -phi, e = e, a = -a, Set = 1)

> E[[2]] < -newTissueFromEllipse(x0 + c(dex, 0), dx = dx, +
```

```
phi, e, a, Set = 2)
> E[[3]] \leftarrow newTissueFromEllipse(x0 + c(-dey, dey), dx = dx,
      -phi, e, -a, Set = 3)
> E[[4]] \leftarrow newTissueFromEllipse(x0 + c(dex + dey, dey),
      dx = dx, phi, e, a, Set = 4)
> TM <- E[[1]]
> TM <- addSetToDrawing(TM, E[[2]], set2Name = "Set2")
> grid.newpage()
> pushViewport(plotViewport(c(1, 1, 1, 1)))
> makevp.eqsc(c(-10, 10), c(-10, 10))
> grid.xaxis()
> grid.yaxis()
> PlotFaces(TM)
> PlotSetBoundaries(TM, gp = gpar(lwd = 2, col = c("green",
      "red", "blue")))
> PlotNodes(TM)
> .PlotFaceNames.TissueDrawing(TM)
```

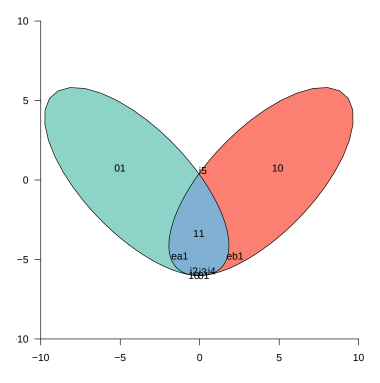


Figure 7: Injecting points

19 This document

Author	Jonathan Swinton
CVS id of this document	Id: TissueDrawingTest.Rnw 14 2009-07-16 09:48:12Z js229.
Generated on	17 th July, 2009
R version	