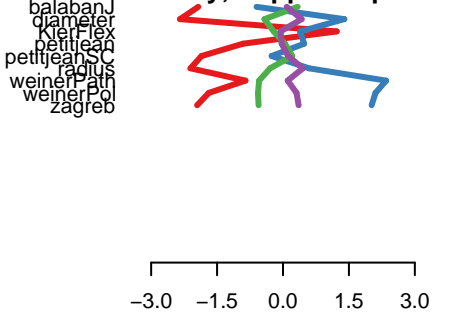
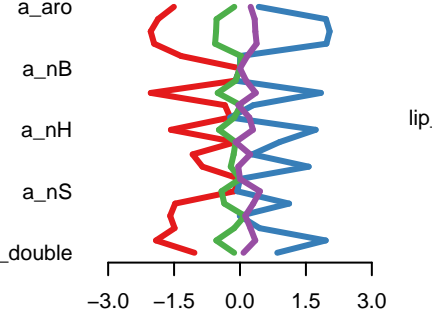


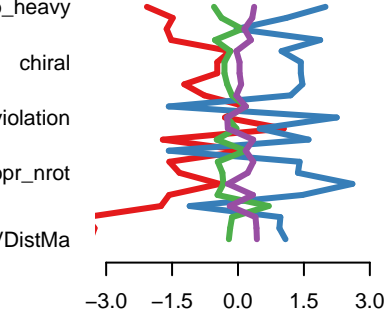
(EII,4,6013) Adjacency, distance matrix, connectivity, Kappa shape in



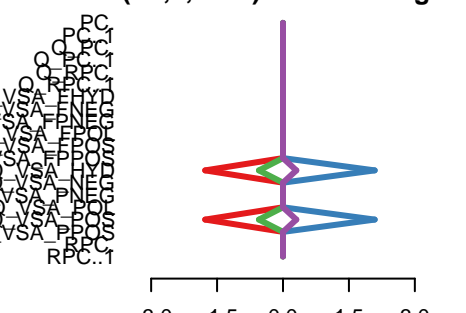
(EII,4,6013) Atom and bond con



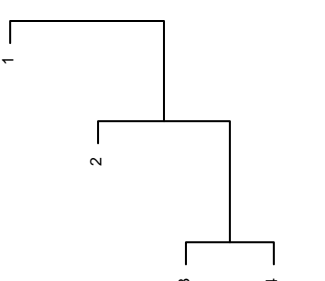
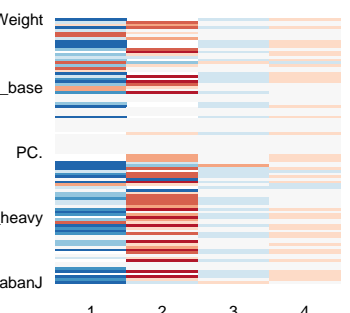
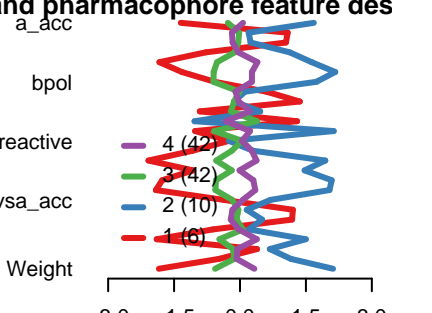
(EII,4,6013) Atom and bond con



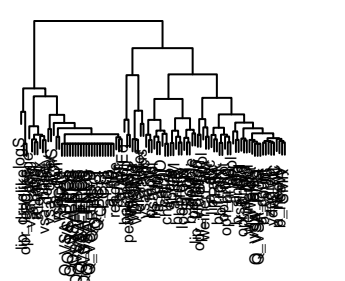
(EII,4,6013) Partial charge



(EII,4,6013) Physical proper and pharmacophore feature des



dist(avg_p)
hclust (*, "complete")



dist(t(avg_p))
hclust (*, "complete")