

test manhattan, umichigan version

Based on https://genome.sph.umich.edu/wiki/Code_Sample:_Generating_Manhattan_Plots_in_R

```
library(lattice)
manhattan.plot<-function(chr, pos, pvalue,
  sig.level=NA, annotate=NULL, ann.default=list(),
  should.thin=T, thin.pos.places=2, thin.logp.places=2,
  xlab="Chromosome", ylab=expression(-log[10](p-value)),
  col=c("gray","darkgray"), panel.extra=NULL, pch=20, cex=0.8,...) {

  if (length(chr)==0) stop("chromosome vector is empty")
  if (length(pos)==0) stop("position vector is empty")
  if (length(pvalue)==0) stop("pvalue vector is empty")

  #make sure we have an ordered factor
  if(!is.ordered(chr)) {
    chr <- ordered(chr)
  } else {
    chr <- chr[,drop=T]
  }

  #make sure positions are in kbp
  if (any(pos>1e6)) pos<-pos/1e6;

  #calculate absolute genomic position
  #from relative chromosomal positions
  posmin <- tapply(pos,chr, min);
  posmax <- tapply(pos,chr, max);
  posshift <- head(c(0,cumsum(posmax)),-1);
  names(posshift) <- levels(chr)
  genpos <- pos + posshift[chr];
  getGenPos<-function(cchr, cpos) {
    p<-posshift[as.character(cchr)]+cpos
    return(p)
  }

  #parse annotations
  grp <- NULL
  ann.settings <- list()
  label.default<-list(x="peak",y="peak",adj=NULL, pos=3, offset=0.5,
    col=NULL, fontface=NULL, fontsize=NULL, show=F)
  parse.label<-function(rawval, groupname) {
    r<-list(text=groupname)
    if(is.logical(rawval)) {
      if(!rawval) {r$show <- F}
    } else if (is.character(rawval) || is.expression(rawval)) {
      if(nchar(rawval)>=1) {
        r$text <- rawval
      }
    }
  }
}
```

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    }
  } else if (is.list(rawval)) {
    r <- modifyList(r, rawval)
  }
  return(r)
}

if(!is.null(annotate)) {
  if (is.list(annotate)) {
    grp <- annotate[[1]]
  } else {
    grp <- annotate
  }
  if (!is.factor(grp)) {
    grp <- factor(grp)
  }
} else {
  grp <- factor(rep(1, times=length(pvalue)))
}

ann.settings<-vector("list", length(levels(grp)))
ann.settings[[1]]<-list(pch=pch, col=col, cex=cex, fill=col, label=label.default)

if (length(ann.settings)>1) {
  lcols<-trellis.par.get("superpose.symbol")$col
  lfills<-trellis.par.get("superpose.symbol")$fill
  for(i in 2:length(levels(grp))) {
    ann.settings[[i]]<-list(pch=pch,
      col=lcols[(i-2) %% length(lcols) +1 ],
      fill=lfills[(i-2) %% length(lfills) +1 ],
      cex=cex, label=label.default);
    ann.settings[[i]]$label$show <- T
  }
  names(ann.settings)<-levels(grp)
}
for(i in 1:length(ann.settings)) {
  if (i>1) {ann.settings[[i]] <- modifyList(ann.settings[[i]], ann.default)}
  ann.settings[[i]]$label <- modifyList(ann.settings[[i]]$label,
    parse.label(ann.settings[[i]]$label, levels(grp)[i]))
}
if(is.list(annotate) && length(annotate)>1) {
  user.cols <- 2:length(annotate)
  ann.cols <- c()
  if(!is.null(names(annotate[-1])) && all(names(annotate[-1])!="")) {
    ann.cols<-match(names(annotate)[-1], names(ann.settings))
  } else {
    ann.cols<-user.cols-1
  }
  for(i in seq_along(user.cols)) {
    if(!is.null(annotate[[user.cols[i]]]$label)) {
      annotate[[user.cols[i]]]$label<-parse.label(annotate[[user.cols[i]]]$label,
        levels(grp)[ann.cols[i]])
    }
  }
}

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        ann.settings[[ann.cols[i]]]<-modifyList(ann.settings[[ann.cols[i]]],
        annotate[[user.cols[i]]])
    }
}
rm(annotate)

#reduce number of points plotted
if(should.thin) {
    thinned <- unique(data.frame(
        logp=round(-log10(pvalue),thin.logp.places),
        pos=round(genpos,thin.pos.places),
        chr=chr,
        grp=grp)
    )
    logp <- thinned$logp
    genpos <- thinned$pos
    chr <- thinned$chr
    grp <- thinned$grp
    rm(thinned)
} else {
    logp <- -log10(pvalue)
}
rm(pos, pvalue)
gc()

#custom axis to print chromosome names
axis.chr <- function(side,...) {
    if(side=="bottom") {
        panel.axis(side=side, outside=T,
            at=((posmax+posmin)/2+posshift),
            labels=levels(chr),
            ticks=F, rot=0,
            check.overlap=F
        )
    } else if (side=="top" || side=="right") {
        panel.axis(side=side, draw.labels=F, ticks=F);
    }
    else {
        axis.default(side=side,...);
    }
}

#make sure the y-lim covers the range (plus a bit more to look nice)
prepanel.chr<-function(x,y,...) {
    A<-list();
    maxy<-ceiling(max(y, ifelse(!is.na(sig.level), -log10(sig.level), 0)))+.5;
    A$ylim=c(0,maxy);
    A;
}

xyplot(logp~genpos, chr=chr, groups=grp,
    axis=axis.chr, ann.settings=ann.settings,
    prepanel=prepanel.chr, scales=list(axes="i"),

```

```

panel=function(x, y, ..., getgenpos) {
  if(!is.na(sig.level)) {
    #add significance line (if requested)
    panel.abline(h=-log10(sig.level), lty=2);
  }
  panel.superpose(x, y, ..., getgenpos=getgenpos);
  if(!is.null(panel.extra)) {
    panel.extra(x,y, getgenpos, ...)
  }
},
panel.groups = function(x,y,..., subscripts, group.number) {
  A<-list(...)
  #allow for different annotation settings
  gs <- ann.settings[[group.number]]
  A$col.symbol <- gs$col[(as.numeric(chr[subscripts])-1) %% length(gs$col) + 1]
  A$cex <- gs$cex[(as.numeric(chr[subscripts])-1) %% length(gs$cex) + 1]
  A$pch <- gs$pch[(as.numeric(chr[subscripts])-1) %% length(gs$pch) + 1]
  A$fill <- gs$fill[(as.numeric(chr[subscripts])-1) %% length(gs$fill) + 1]
  A$x <- x
  A$y <- y
  do.call("panel.xyplot", A)
  #draw labels (if requested)
  if(gs$label$show) {
    gt<-gs$label
    names(gt)[which(names(gt)=="text")]<-"labels"
    gt$show<-NULL
    if(is.character(gt$x) | is.character(gt$y)) {
      peak = which.max(y)
      center = mean(range(x))
      if (is.character(gt$x)) {
        if(gt$x=="peak") {gt$x<-x[peak]}
        if(gt$x=="center") {gt$x<-center}
      }
      if (is.character(gt$y)) {
        if(gt$y=="peak") {gt$y<-y[peak]}
      }
    }
    if(is.list(gt$x)) {
      gt$x<-A$getgenpos(gt$x[[1]],gt$x[[2]])
    }
    do.call("panel.text", gt)
  }
},
xlab=xlab, ylab=ylab,
panel.extra=panel.extra, getgenpos=getGenPos, ...
);
}

```

```
list.files()
```

```

## [1] "gwas-t2m-meta0613-positions-pq.csv" "michigan.Rmd"
## [3] "msa_position_2_ref_position.csv"    "plot_gisaid_gwas-t2m.pdf"
## [5] "plot_gisaid_gwas-t2m.Rmd"          "README.md"

```

```

tb = read.csv2( "gwas-t2m-meta0613-positions-pq.csv", stringsAsFactors = F)
#tb

tb_cov = read.csv("msa_position_2_ref_position.csv", row.names = 1)
tb$p = as.numeric(tb$p)
tb$ref_position = tb_cov$ref_positions[ match(tb$msa_position, tb_cov$msa_positions )]
tb$p[tb$p==0] = 1E-200
summary(tb)

##      R_position      msa_position      p      q
## Length:4210      Min.   : 582      Min.   :0.0000000      Length:4210
## Class :character      1st Qu.: 8827      1st Qu.:0.0000000      Class :character
## Mode  :character      Median :28652      Median :0.0000039      Mode  :character
##                               Mean  :23641      Mean   :0.0789487
##                               3rd Qu.:34110      3rd Qu.:0.0581378
##                               Max.   :48402      Max.   :0.9983663
##      ref_position
##      Min.   : 0
##      1st Qu.: 5500
##      Median :19488
##      Mean   :15095
##      3rd Qu.:21489
##      Max.   :29890
tb$chr=1

# spike 21563 25384

#make annotation factor
ann <- rep(1, length(tb$p))
#print(ann)
ann[with(tb, chr==1 & ref_position>=21563 & ref_position< 25384)]<-2
#print(ann)
ann<-factor(ann, levels=1:2, labels=c("", "spike"))

#draw plot with annotation

#manhattan.plot(tb$chr, tb$ref_position, tb$p)
#draw plot with annotation
manhattan.plot(tb$chr, tb$ref_position, tb$p, annotate=ann)

```

