ggbeeswarm package usage example (version 0.6.0)

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Abstract

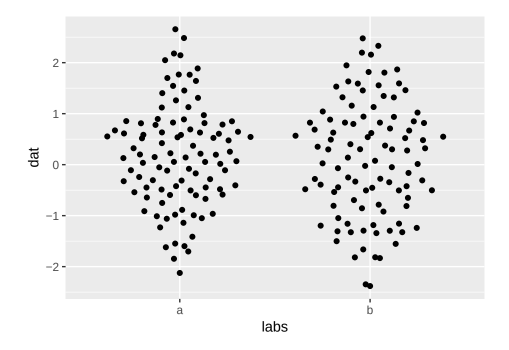
This is a collection of examples of usage for the **ggbeeswarm** package.

Keywords: visualization, display, one dimensional, grouped, groups, violin, scatter, points, quasirandom, beeswarm, van der Corput, beeswarm, ggplot, ggplot2.

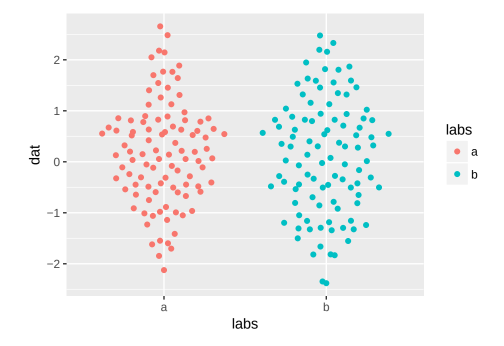
1. The basics

This is the simplest example of using geom_quasirandom to generate violin scatter plots:

- > library(ggbeeswarm)
- > set.seed(12345)
- > n<-100
- > dat<-rnorm(n*2)</pre>
- > labs<-rep(c('a','b'),n)</pre>
- > ggplot(mapping=aes(labs, dat)) + geom_quasirandom()



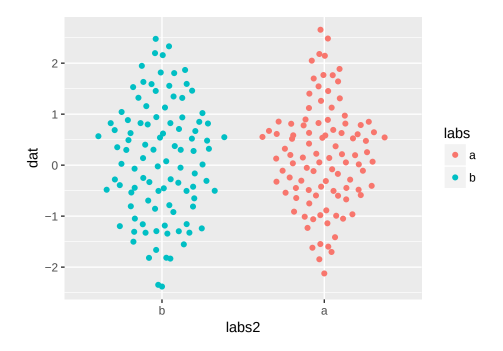
Normal ggplot options can be used:



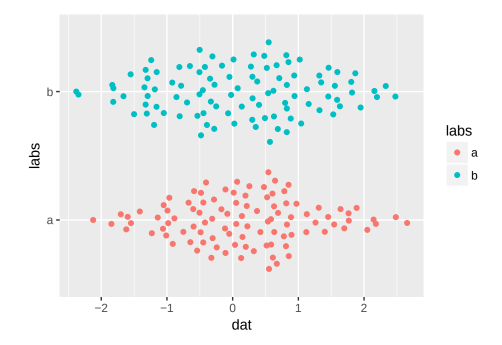
> ggplot(mapping=aes(labs, dat)) + geom_quasirandom(aes(color=labs))

Factors can be used to generate custom group orderings:

- > labs2<-factor(labs,levels=c('b','a'))</pre>
- > ggplot(mapping=aes(labs2, dat)) + geom_quasirandom(aes(color=labs))



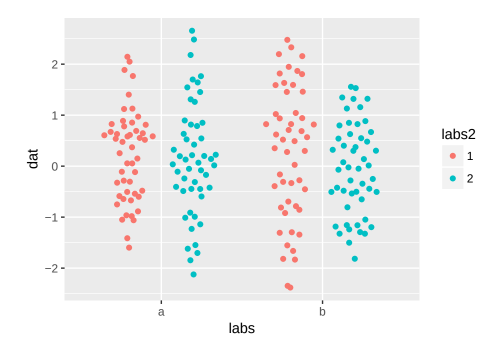
The axes can also be switched with a categorical y-axis using the argument groupOnX=FALSE:



> ggplot(mapping=aes(dat,labs)) + geom_quasirandom(aes(color=labs),groupOnX=FALSE)

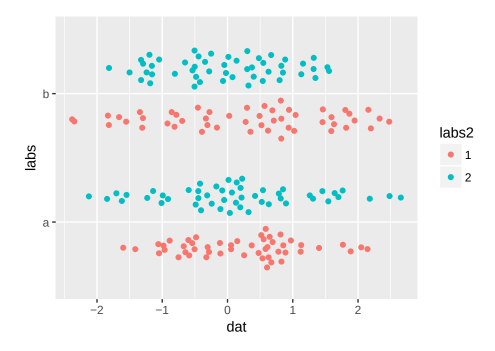
And dodging can be used to compare within groups:

- > labs2<-factor(rep(1:2,each=n))</pre>
- > ggplot(mapping=aes(labs,dat,color=labs2)) + geom_quasirandom(dodge.width=.8)



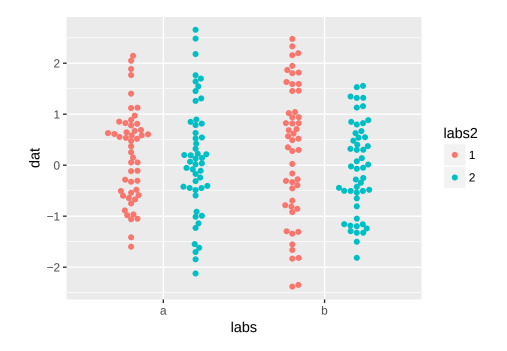
Or on the y-axis:

- > labs2<-factor(rep(1:2,each=n))</pre>
- > ggplot(mapping=aes(dat,labs,color=labs2)) + geom_quasirandom(dodge.width=.8,groupOnX=F



And with geom_beeswarm:

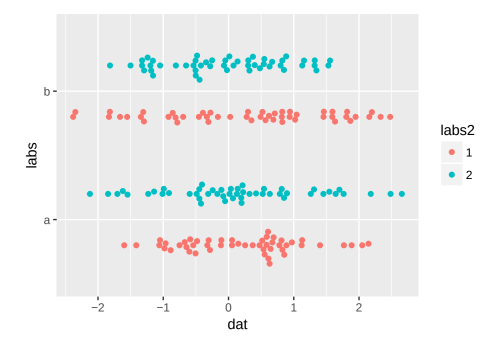
- > ggplot(mapping=aes(labs,dat,color=labs2)) +
- + geom_beeswarm(dodge.width=.8,cex=2)



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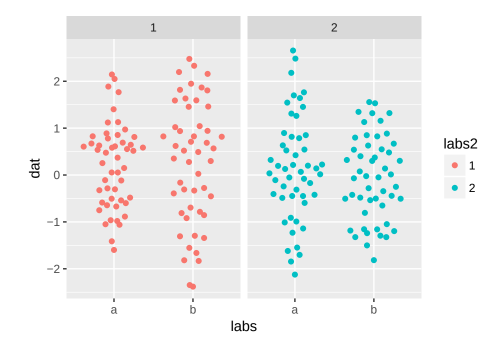
```
> ggplot(mapping=aes(dat,labs,color=labs2)) +
```

+ geom_beeswarm(dodge.width=.8,cex=2,groupOnX=FALSE)



Both geom_beeswarm and geom_quasirandom also work with facets:

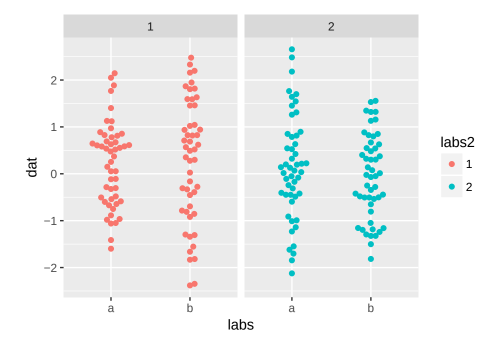
- > df<-data.frame(labs,dat,labs2)
- > ggplot(df,aes(labs,dat,color=labs2)) +
- + geom_quasirandom() +
- + facet_grid(.~labs2)



```
> ggplot(df,aes(labs,dat,color=labs2)) +
```

```
+ geom_beeswarm(cex=3) +
```

```
+ facet_grid(.~labs2)
```





There are several ways to plot grouped one-dimensional data combining points and density estimation including:

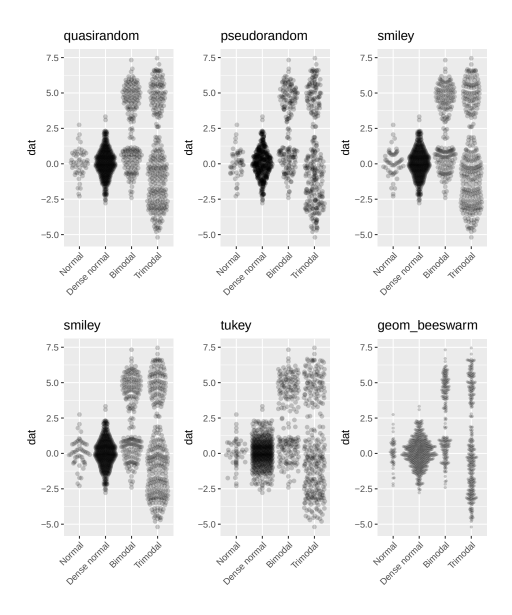
- **pseudorandom** The kernel density is estimated then points are distributed uniform randomly within the density estimate for a given bin. Selection of an appropriate number of bins does not greatly affect appearance but coincidental clumpiness is common.
- alternating within bins The kernel density is estimated then points are distributed within the density estimate for a given bin evenly spaced with extreme values alternating from right to left e.g. max, 3rd max, ..., 4th max, 2nd max. If maximums are placed on the outside then these plots often form consecutive "smiley" patterns. If minimums are placed on the outside then "frowny" patterns are generated. Selection of the number of bins can have large effects on appearance important.
- tukey An algorithm described by Tukey and Tukey in "Strips displaying empirical distributions: I. textured dot strips" using constrained permutations of offsets to distrbute the data.
- **quasirandom** The kernel density is estimated then points are distributed quasirandomly using the von der Corput sequence within the density estimate for a given bin. Selection of an appropriate number of bins does not greatly affect appearance and position does not depend on plotting parameters.

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beeswarm The package **beeswarm** provides methods for generating a "beeswarm" plot where points are distibuted so that no points overlap. Kernel density is not calculated although the resulting plot does provide an approximate density estimate. Selection of an appropriate number of bins affects appearance and plot and point sizes must be known in advance.

The first four options are included within geom_quasirandom using the method= argument and beeswarm plots are generated with geom_beeswarm:

```
>
    library(gridExtra)
>
    dat <- list(</pre>
      'Normal'=rnorm(50),
+
      'Dense normal'= rnorm(500),
+
      'Bimodal'=c(rnorm(100), rnorm(100,5)),
      'Trimodal'=c(rnorm(100), rnorm(100,5),rnorm(100,-3))
+
    )
+
    labs<-rep(names(dat),sapply(dat,length))</pre>
>
    labs<-factor(labs,levels=unique(labs))</pre>
>
    dat<-unlist(dat)</pre>
>
    p1<-ggplot(mapping=aes(labs, dat)) +</pre>
>
      geom_quasirandom(alpha=.2) +
+
+
      ggtitle('quasirandom') + labs(x='') +
      theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1))
+
    p2<-ggplot(mapping=aes(labs, dat)) +</pre>
>
+
      geom_quasirandom(method='pseudorandom',alpha=.2) +
+
      ggtitle('pseudorandom') + labs(x='') +
      theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1))
+
    p3<-ggplot(mapping=aes(labs, dat)) +</pre>
>
+
      geom_quasirandom(method='smiley',alpha=.2) +
+
      ggtitle('smiley') + labs(x='') +
      theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1))
+
>
    p4<-ggplot(mapping=aes(labs, dat)) +</pre>
      geom_quasirandom(method='frowney',alpha=.2) +
+
      ggtitle('smiley') + labs(x='') +
+
      theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1))
+
    p5<-ggplot(mapping=aes(labs, dat)) +</pre>
>
      geom_quasirandom(method='tukey',alpha=.2) +
+
      ggtitle('tukey') + labs(x='') +
+
      theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1))
+
>
    p6<-ggplot(mapping=aes(labs, dat)) +</pre>
+
      geom_beeswarm(alpha=.2,size=.75) +
+
      ggtitle('geom_beeswarm') + labs(x='') +
      theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust=1))
+
>
    grid.arrange(p1, p2, p3, p4, p5, p6, ncol=3)
```

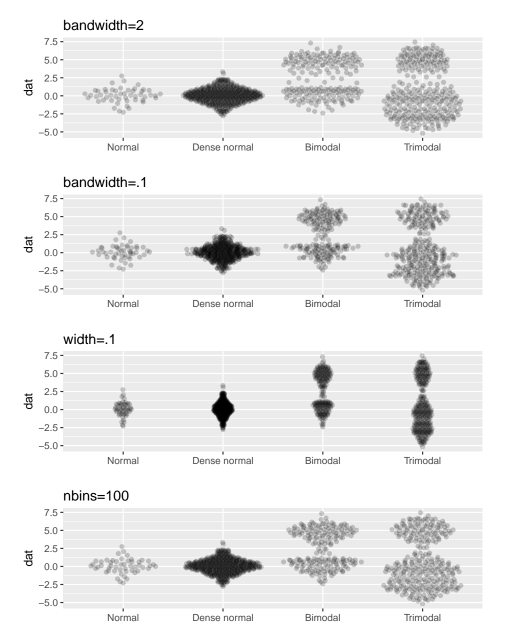


quasirandom calls vipor::offsetX which calls stats::density to compute kernel density estimates. The tightness of the fit can be adjusted with the bandwidth option and the width of the offset with width. nbins to adjust the number of bins used in the kernel density is also provided but this can usually be left at its default when using quasirandom offsets but is useful for non-quasirandom methods:

```
> library(gridExtra)
> p1<-ggplot(mapping=aes(labs, dat)) +
    geom_quasirandom(bandwidth=2,alpha=.2) +
    ggtitle('bandwidth=2') + labs(x='')
> p2<-ggplot(mapping=aes(labs, dat)) +
    geom_quasirandom(bandwidth=.1,alpha=.2) +
    ggtitle('bandwidth=.1') + labs(x='')
```

> p3<-ggplot(mapping=aes(labs, dat)) +</pre>

```
+ geom_quasirandom(width=.1,alpha=.2) +
    ggtitle('width=.1') + labs(x='')
> p4<-ggplot(mapping=aes(labs, dat)) +
    geom_quasirandom(nbins=100,alpha=.2) +
    ggtitle('nbins=100') + labs(x='')
> grid.arrange(p1, p2, p3, p4, ncol=1)
```



The frowney or smiley methods are sensitive to the number of bins so the argument nbins is more useful/necessary with them:

```
> p1<-ggplot(mapping=aes(labs, dat)) +</pre>
```

```
+ geom_quasirandom(method='smiley',alpha=.2) +
```

```
+ ggtitle('Default (n/5)') + labs(x='')
```

> p2<-ggplot(mapping=aes(labs, dat)) +</pre>

```
+ geom_quasirandom(method='smiley',nbins=50,alpha=.2) +
```

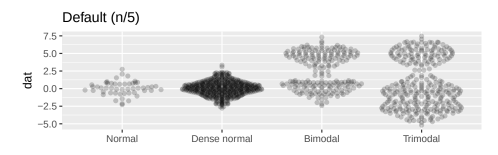
- + ggtitle('nbins=50') + labs(x='')
- > p3<-ggplot(mapping=aes(labs, dat)) +</pre>

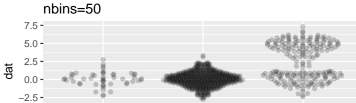
```
+ geom_quasirandom(method='smiley',nbins=100,alpha=.2) +
```

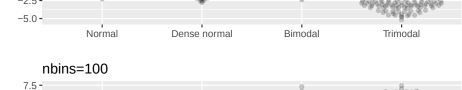
```
+ ggtitle('nbins=100') + labs(x='')
```

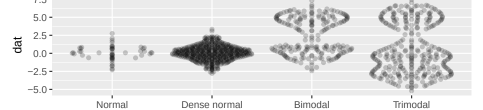
```
> p4<-ggplot(mapping=aes(labs, dat)) +</pre>
```

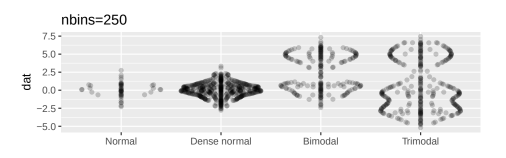
- + geom_quasirandom(method='smiley',nbins=250,alpha=.2) +
- + ggtitle('nbins=250') + labs(x='')
- > grid.arrange(p1, p2, p3, p4, ncol=1)







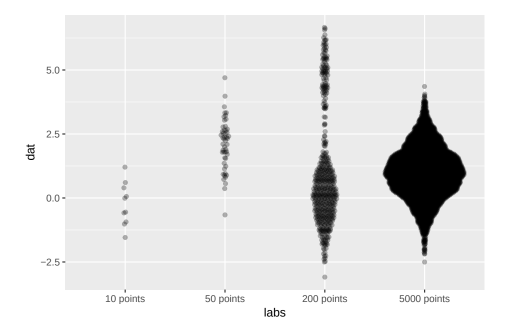


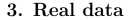


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The varwidth argument scales the width of a group by the square root of the number of observations in that group (as in the function boxplot):

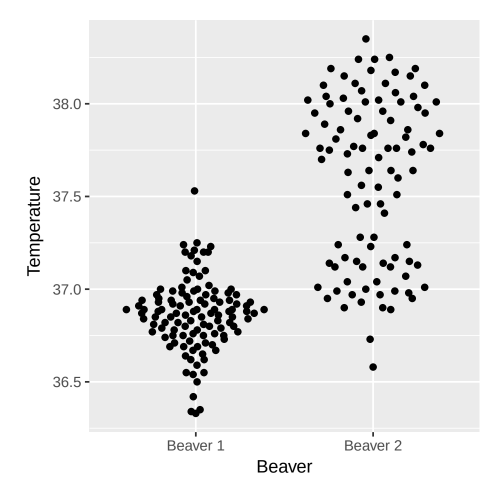
```
dat <- list(</pre>
>
+
       '10 points'=rnorm(10),
       '50 points'=rnorm(50,2),
+
       '200 points'=c(rnorm(400), rnorm(100,5)),
+
+
       '5000 points'= rnorm(5000,1)
    )
+
>
    labs<-rep(names(dat),sapply(dat,length))</pre>
    labs<-factor(labs,levels=unique(labs))</pre>
>
>
    dat<-unlist(dat)</pre>
    ggplot(mapping=aes(labs, dat)) + geom_quasirandom(alpha=.3,varwidth=TRUE)
>
```





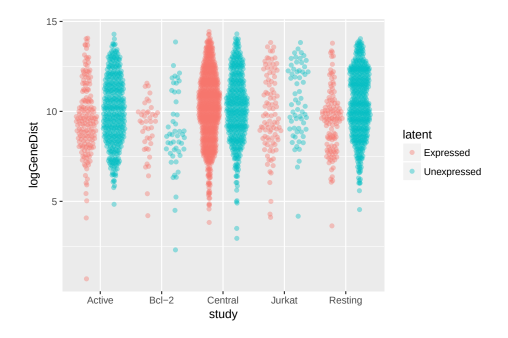
An example using the beaver1 and beaver2 data from the datasets package:

```
>
    beaver<-data.frame(</pre>
      'Temperature'=c(beaver1$temp,beaver2$temp),
+
      'Beaver'=rep(
+
        c('Beaver 1', 'Beaver 2'),
+
        c(nrow(beaver1),nrow(beaver2))
+
      )
+
+
    )
>
    ggplot(beaver,mapping=aes(Beaver, Temperature)) + geom_quasirandom()
```



An example using the integrations data from the vipor package and the argument dodge.width:

- > library(vipor)
- > ints<-integrations[integrations\$nearestGene>0,]
- > ints\$logGeneDist<-log(ints\$nearestGene)</pre>
- > ggplot(ints,mapping=aes(study, logGeneDist,color=latent)) +
- + geom_quasirandom(dodge.width=.9,alpha=.4)



Affiliation:

Github: http://github.com/eclarke/ggbeeswarm Cran: https://cran.r-project.org/package=ggbeeswarm