# Statistical Inference: matched pairs and normal quantile plot

# Matched pairs

#### Some data:

	_	
subject	druga	drugb
1	2.0	3.5
2	3.6	5.7
3	2.6	2.9
4	2.6	2.4
5	7.3	9.9
6	3.4	3.3
1 2 3 4 5 6 7 8 9	14.9	16.7
8	6.6	6.0
9	2.3	3.8
10	2.0	4.0
11	6.8	9.1
12	8.5	20.9

# Matched pairs data

- Data are comparison of 2 drugs for effectiveness at reducing pain.
  - 12 subjects (cases) were arthritis sufferers
  - Response is #hours of pain relief from each drug.
- In reading example, each child tried only one reading method.
- But here, each subject tried out both drugs, giving us two measurements.
  - Possible because, if you wait long enough, one drug has no influence over effect of other.
  - Advantage: focused comparison of drugs. Compare one drug with another on same person, removes a lot of variability due to differences between people.
  - Matched pairs, requires different analysis.
- Design: randomly choose 6 of 12 subjects to get drug A first, other 6 get drug B first.

## Paired t test: reading the data

subject = col\_double(),
druga = col\_double(),

drugb = col double()

```
Values aligned in columns:
```

##

##

## ## )

```
my_url="http://www.utsc.utoronto.ca/~butler/c32/analgesic.txt"
pain=read_table(my_url)
## Parsed with column specification:
## cols(
```

## The data

pain

subject	druga	drugb
1	2.0	3.5
2	3.6	5.7
3	2.6	2.9
4	2.6	2.4
5	7.3	9.9
6	3.4	3.3
7	14.9	16.7
8	6.6	6.0
9	2.3	3.8
10	2.0	4.0
11	6.8	9.1
12	8.5	20.9

#### Paired t-test

```
with(pain, t.test(druga, drugb, paired = T))
##
##
   Paired t-test
##
## data: druga and drugb
## t = -2.1677, df = 11, p-value = 0.05299
## alternative hypothesis: true difference in means is not equal to
## 95 percent confidence interval:
## -4.29941513 0.03274847
## sample estimates:
## mean of the differences
##
                 -2.133333
```

- P-value is 0.053
- Not quite evidence of difference between drugs.

## t-testing the differences

- Likewise, you can calculate the differences yourself and do a 1-sample t-test on them.
- First calculate a column of differences:

(pain %>% mutate(diff=druga-drugb) -> pain)

subject	druga	drugb	diff
1	2.0	3.5	-1.5
2	3.6	5.7	-2.1
3	2.6	2.9	-0.3
4	2.6	2.4	0.2
5	7.3	9.9	-2.6
6	3.4	3.3	0.1
7	14.9	16.7	-1.8
8	6.6	6.0	0.6
9	2.3	3.8	-1.5
10	2.0	4.0	-2.0
11	6.8	9.1	-2.3
12	8.5	20.9	-12.4

#### t-test on the differences

## -2.133333

• then throw them into t.test, testing that the mean is zero, with same result as before:

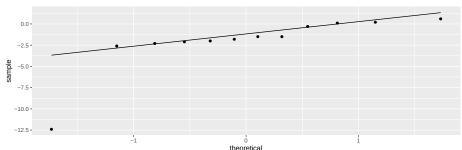
```
with(pain,t.test(diff,mu=0))
##
   One Sample t-test
##
##
## data: diff
## t = -2.1677, df = 11, p-value = 0.05299
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
## -4.29941513 0.03274847
## sample estimates:
## mean of x
```

# Assessing normality

- 1-sample and 2-sample t-tests assume (each) group normally distributed.
- Matched pairs analyses assume (theoretically) that differences normally distributed.
- Though we know that t-tests generally behave well even without normality.
- How to assess normality? A normal quantile plot.
  - Idea: scatter of points should follow the straight line, without curving.
  - Outliers show up at bottom left or top right of plot as points off the line.

# The normal quantile plot

of differences from matched pairs data



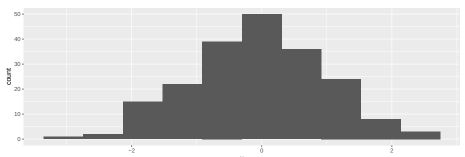
 Points should follow the straight line. Bottom left one way off, so normality questionable here: outlier.

# More normal quantile plots

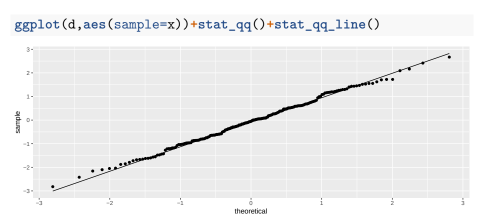
- How straight does a normal quantile plot have to be?
- There is randomness in real data, so even a normal quantile plot from normal data won't look perfectly straight.
- With a small sample, can look not very straight even from normal data.
- Looking for systematic departure from a straight line; random wiggles ought not to concern us.
- Look at some examples where we know the answer, so that we can see what to expect.

## Normal data, large sample





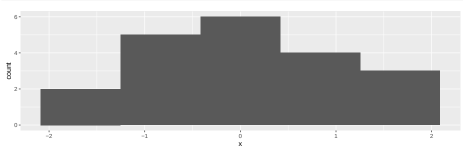
# The normal quantile plot



## Normal data, small sample

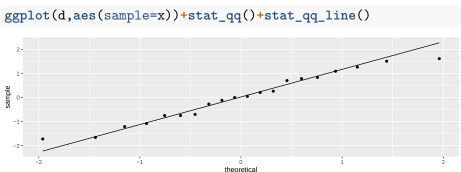
Not so convincingly normal, but not obviously skewed:

```
d=tibble(x=rnorm(20))
ggplot(d,aes(x=x))+geom_histogram(bins=5)
```



# The normal quantile plot

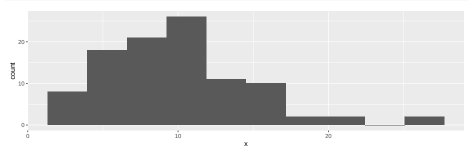
Good, apart from the highest and lowest points being slightly off. I'd call this good:



## Chi-squared data, df = 10

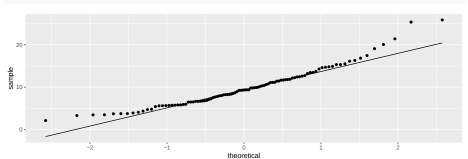
#### Somewhat skewed to right:

```
d=tibble(x=rchisq(100,10))
ggplot(d,aes(x=x))+geom_histogram(bins=10)
```



# The normal quantile plot

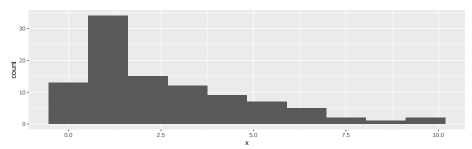
#### Somewhat opening-up curve:



## Chi-squared data, df = 3

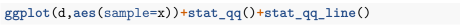
#### Definitely skewed to right:

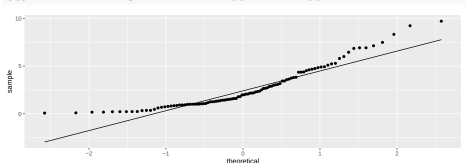
```
d=tibble(x=rchisq(100,3))
ggplot(d,aes(x=x))+geom_histogram(bins=10)
```



# The normal quantile plot

#### Clear upward-opening curve:

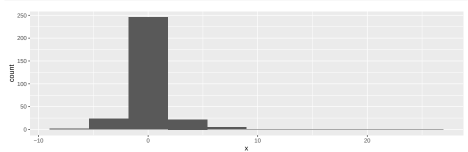




## t-distributed data, df = 3

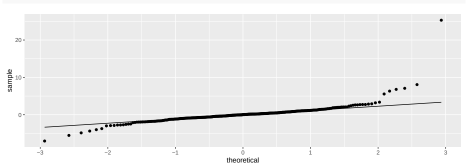
Long tails (or a very sharp peak):

```
d=tibble(x=rt(300,3))
ggplot(d,aes(x=x))+geom_histogram(bins=10)
```

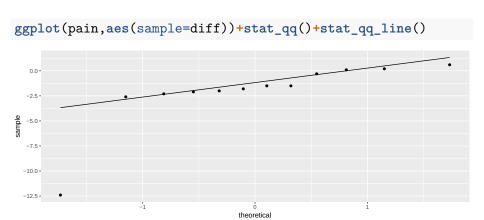


# The normal quantile plot

Low values too low and high values too high for normal.



# Our pain-relief data



## Comments

- Definitely not normal. What to do?
- Sign test on differences, null median 0.

# Sign test

- Most easily: calculate differences in data frame, then use smmr.
- Null median difference is 0:

```
pain %>% mutate(mydiff=druga-drugb) %>%
sign_test(mydiff,0)
```

```
## $above_below
## below above
## 9 3
##
## $p_values
## alternative p_value
## 1 lower 0.07299805
## 2 upper 0.98071289
## 3 two-sided 0.14599609
```

#### Comments

- P-value 0.1460. No evidence that the drugs are different.
- Since we are working in a pipeline, input data frame to sign\_test is "whatever came out of previous step".

# (Some of) the kids' reading data, again

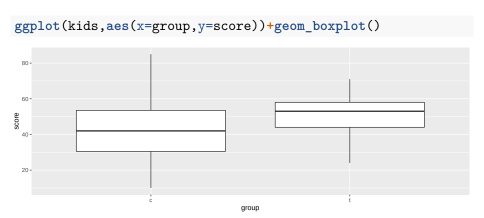
kids %>% slice\_sample(n=12)

group	score
t	49
С	48
С	46
С	17
t	43
t	44
С	26
t	59
С	55
t	62
t	43
С	10

## Where we were at

- 21 kids in "treatment", new reading method; 23 in "control", standard reading method.
- Assessing assumptions:
  - We did two-sample t-test (Satterthwaite-Welch) before.
  - Assumes approx. normal data within each group.
  - Does not assume equal spread.
  - (Pooled t-test *does* assume equal spread).
  - Assess each group separately.

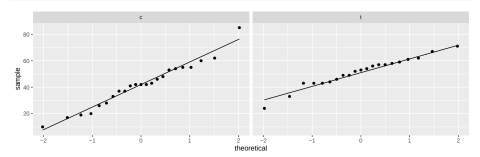
## Boxplots for reading data



## Facetted normal quantile plots

#### Done this way:

```
ggplot(kids,aes(sample=score))+stat_qq()+stat_qq_line()+
facet_wrap(~group)
```



## Comments

- These plots show no problems with normality. Both groups are more or less symmetric/normal and there are no outliers.
- Equal spreads questionable, but we don't need that.
- Assess equal spreads by looking at *slopes* of normal quantile plots.
- We ought be happy with the (Welch) two-sample t-test (over)

# Welch two-sample test

##

```
t.test(score~group,data=kids,alternative="less")
##
##
   Welch Two Sample t-test
##
## data: score by group
## t = -2.3109, df = 37.855, p-value = 0.01319
## alternative hypothesis: true difference in means is less tl
## 95 percent confidence interval:
       -Inf -2.691293
##
## sample estimates:
## mean in group c mean in group t
```

from which we concluded that the new reading method really does help.

41.52174 51.47619