

# The cbcTools Package: Tools for Designing and Testing Choice-Based Conjoint Surveys in R

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## Abstract

While many open source packages and applications exist for estimating choice models using data from choice-based conjoint experiments, few packages exist for generating and evaluating experiment designs prior to collecting data. Furthermore, existing tools for designing choice-based conjoint survey experiments often focus on optimizing the design for statistical power under ideal conditions, but they rarely provide guidance on important design decisions for less ideal conditions, such as when preference heterogeneity or strong interactions between certain attributes may be expected in respondent choices. The `cbcTools` R package was developed to provide researchers tools for creating and evaluating experiment designs and sample size requirements under a variety of different conditions prior to fielding an experiment. The package contains functions for generating experiment designs, examining attribute balance and overlap, simulating choice data, and conducting power analyses. The package data format matches that of experiment designs exported from Sawtooth Software, enabling a smooth integration with the Sawtooth Software workflow. Detailed package documentation can be found at <https://jhelvy.github.io/cbcTools/>.

## Introduction

Designing a choice-based conjoint survey is almost never a simple, straightforward process. Designers must consider multiple trade offs between design parameters (e.g., which attributes and levels to include, how many choice questions to ask each respondent, and how many alternatives per choice question) and the design outcomes in terms of the user experience and the statistical power available for identifying effects. The process is typically highly iterative (R. M. Johnson and Orme 2002).

As a quick example, consider a simple conjoint experiment about cars with just two attributes with three levels each:

- **Price:** \$20,000, 40,000, & 100,000
- **Brand:** GM, BMW, & Ferrari

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A common starting point is to first generate the full factorial set of all possible combinations of brand and price. These profiles can then be selected into a survey design from which respondents will make choices. Without any prior knowledge or experience about car prices and brands, a designer may simply choose to create a survey design by randomly selecting profiles from the full factorial set of profiles. This often results in a less efficient design, but it is a starting point for illustrating trade offs in making a design.

Once a survey design is created, one of the first things designers examine is the count of how often each level of each attribute is shown, or the design “balance” (Hauser and Toubia 2005; Huber and Zwerina 1996). The table below shows an example of the counts from a design with just 9 choice sets of 3 alternatives per question, each of which were randomly chosen from the full factorial set of profiles:

Table 1: Individual and pairwise counts of attributes.

Price:		\$20,000	\$40,000	\$100,000
Brand		9	9	9
GM	10	3	0	7
BMW	11	4	5	2
Ferrari	6	2	4	0

Based on the balance alone, it is clear that this design has several problems. First, while the price levels are perfectly balanced (each level is shown 9 times), the brand levels are not – GM and BMW are shown 10 and 11 times, respectively, whereas Ferrari is only shown 6 times. And the pairwise counts are particularly troubling. The Ferrari brand is only shown with a price of \$20,000 and \$40,000 and never at the \$100,000 level (the most logical level for a Ferrari!). Likewise, GM brand is shown with a price of \$100,000 in 7 out of 10 times and never with a price of \$40,000.

This rather poor design is a common outcome when generating a design by randomly selecting profiles from the full factorial with only a limited number of choice sets. One way to improve the outcome is to simply increase the number of choice sets used, which often results in a much better balance (Kuhfeld 2002). For example, if we increase the number of choice sets from 9 to 90, we obtain the following counts:

Table 2: Individual and pairwise counts of attributes.

Price:		\$20,000	\$40,000	\$100,000
Brand		91	84	95
GM	92	31	31	30
BMW	80	25	25	30
Ferrari	98	35	30	35

This design has a much better balance across the attribute levels than the one created from

just 9 choice sets, but it too has issues. Consider, for example, that about 1/3 of the time the Ferrari brand is shown it has a price of just \$20,000. This is obviously an unrealistic scenario, and if a user saw such a profile multiple times they may not take the choice exercise seriously.

As this simple example illustrates, randomly selecting profiles (even with a larger number of choice sets) can still produce a flawed design<sup>1</sup>. An alternative approach is to attempt to identify a statistically optimal design, which is now a rather well-established practice in the literature on discrete choice experiments (F. R. Johnson et al. 2013). Optimal designs maximize the expected Fisher information, which depends on the parameters of the assumed choice model. As a result, a design’s efficiency is related to how accurately the prior guess is to the true parameters. Using a “Bayesian” D-efficient design where the designer can specify prior expected coefficients for each attribute has been proven to outperform other design approaches when sufficient information on the utility model is known (Kessels, Goos, and Vandebroek 2006; Walker et al. 2018).

Table 3: Example conjoint attributes and levels.

Attribute	Level	Prior
Price	\$20,000	0
	\$40,000	-1
	\$100,000	-4
Brand	GM	0
	BMW	1
	Ferrari	2

Using a Bayesian D-efficient design with 90 choice sets and the prior utilities in Table 3, the counts become much more reasonable (see Table 4). Now the individual attribute counts are relatively balanced, and the pairwise combinations of attributes are much more plausible. For example, the GM brand is only shown at price levels of \$20,000 and \$40,000, and about half of the time Ferrari is shown at the \$100,000 price level.

Table 4: Individual and pairwise counts of attributes.

Price:		\$20,000	\$40,000	\$100,000
Brand		97	93	78
GM	93	52	41	0
BMW	90	30	30	30
Ferrari	86	15	22	49

Of course, even Bayesian D-efficient designs have their downsides. In particular, they can

<sup>1</sup>Note that randomly selecting profiles is different from the kinds of “randomized” designs Sawtooth Software generates, where respondents are randomly assigned to blocks in the experiment which later are *not* randomly constructed.

become problematic if there are significant interaction effects between any of the attributes that the designer did not anticipate.

Figure 1 highlights this trade off. The plots show the standard errors of each coefficients from the same model estimated using increasing subsets of two simulated data sets: a randomized design and a Bayesian D-efficient design. In the first row, it is clear that the D-efficient design produces lower standard errors for the same sample size compared to the randomized design. However, the second row shows the standard errors on interaction effects, which are not identifiable using the D-efficient design. This is because interaction effects can become confounded with main effects in D-efficient designs where the designer did not consider interactions in their priors. Thus, the accuracy of the designer’s prior model used in obtaining a D-efficient design is an important factor in the overall performance of the design. If little to no prior information is known and interactions may be present, a fully randomized design may be a better choice, though of course this would require larger sample sizes to obtain significant parameter estimates.

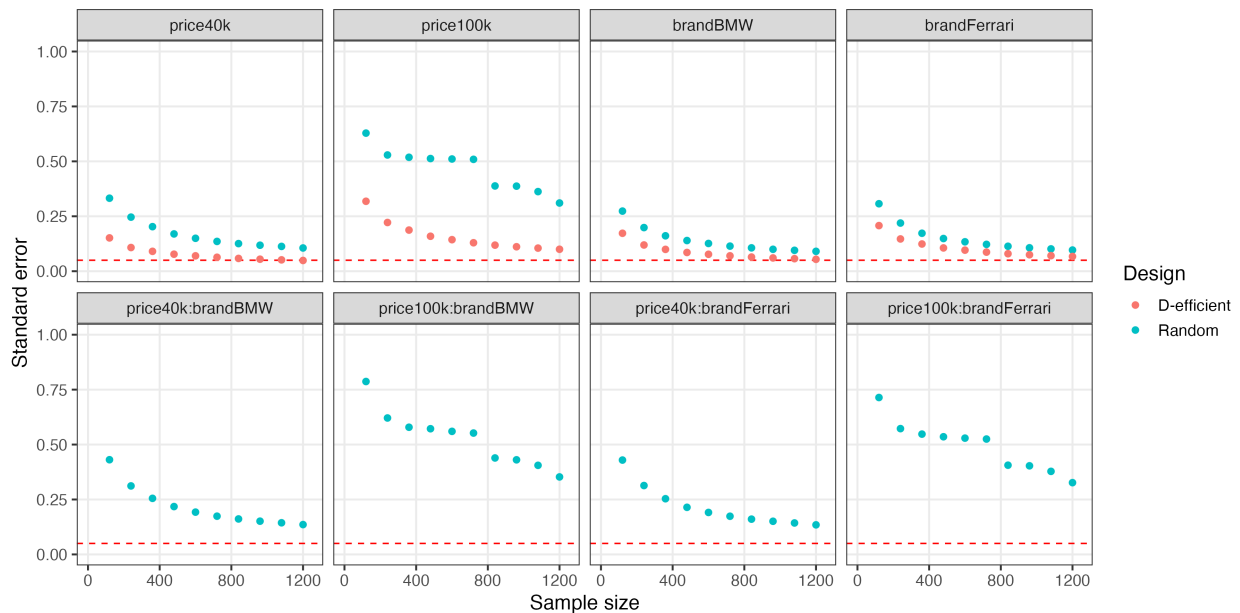


Figure 1: Standard errors with increasing sample size for a multinomial logit model with interaction effects estimated using a randomized design versus a Bayesian D-efficient design

## The cbcTools Package

As the previous example illustrates, designing a choice-based conjoint experiment is an iterative process requiring careful consideration of a variety of factors. The `cbcTools` package was designed as an open source tool to help designers navigate this process and to understand the impacts that different design decisions could have on the outcomes of a choice-based conjoint experiment prior to fielding the survey (Helveston 2022).

The package provides a set of functions (each starting with `cbc_`) for designing surveys and

conducting power analyses for choice-based conjoint survey experiments in R. Each function is associated with a process in the typical design workflow (see Figure 2). The rest of this paper explains each of these steps using a detailed example.

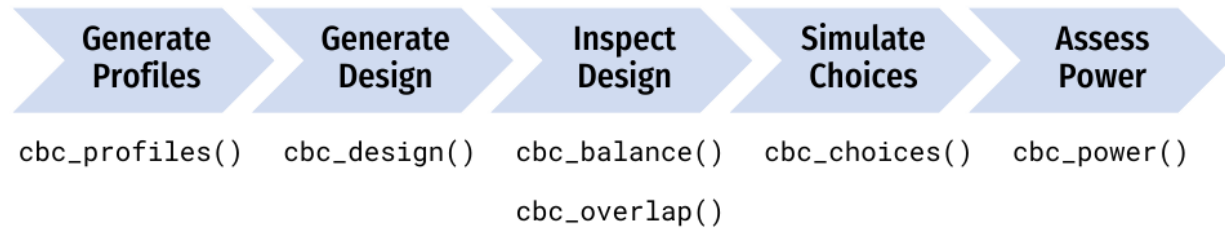


Figure 2: Diagram of the choice-based conjoint survey design process and associated package function(s).

## Generating profiles

One of the first steps in designing a conjoint experiment is to define the attributes and levels and then generate all of the profiles for each combination of those attributes and levels, often referred to as the “full factorial” design. For example, consider designing a conjoint experiment about apples with the following three attributes: `price`, `type`, and `freshness`. The full factorial set of profiles for these attributes can be obtained using the `cbc_profiles()` function:

```
profiles <- cbc_profiles(
  price      = seq(1, 4, 0.5), # $ per pound
  type       = c('Fuji', 'Gala', 'Honeycrisp'),
  freshness  = c('Poor', 'Average', 'Excellent')
)
```

```
head(profiles)
```

```
#>   profileID price type freshness
#> 1         1  1.0 Fuji     Poor
#> 2         2  1.5 Fuji     Poor
#> 3         3  2.0 Fuji     Poor
#> 4         4  2.5 Fuji     Poor
#> 5         5  3.0 Fuji     Poor
#> 6         6  3.5 Fuji     Poor
```

```
tail(profiles)
```

```
#>   profileID price      type freshness
#> 58         58  1.5 Honeycrisp Excellent
#> 59         59  2.0 Honeycrisp Excellent
#> 60         60  2.5 Honeycrisp Excellent
```

```
#> 61      61    3.0 Honeycrisp Excellent
#> 62      62    3.5 Honeycrisp Excellent
#> 63      63    4.0 Honeycrisp Excellent
```

The resulting data frame, `profiles`, has 63 rows, each defining a unique profile. Depending on the context of the survey, some profiles may need to be eliminated (e.g., some profile combinations may be illogical or unrealistic).<sup>2</sup> To do so, each level of an attribute can be defined as a list defining those constraints. In the example below, the `type` attribute has constraints such that only certain price levels will be shown for each level. In addition, for the "Honeycrisp" level of the `type` attribute, only two of the three `freshness` levels are included: "Excellent" and "Average". Note that both the other attributes (`price` and `freshness`) should contain all of the possible levels. With these constraints, only 30 profiles are available compared to 63 without constraints.

```
profiles <- cbc_profiles(
  price = c(1, 1.5, 2, 2.5, 3, 3.5, 4, 4.5, 5),
  freshness = c('Poor', 'Average', 'Excellent'),
  type = list(
    "Fuji" = list(
      price = c(2, 2.5, 3)
    ),
    "Gala" = list(
      price = c(1, 1.5, 2)
    ),
    "Honeycrisp" = list(
      price = c(2.5, 3, 3.5, 4, 4.5, 5),
      freshness = c("Average", "Excellent")
    )
  )
)

head(profiles)
```

```
#>  profileID price freshness type
#> 1         1  2.0      Poor Fuji
#> 2         2  2.5      Poor Fuji
#> 3         3  3.0      Poor Fuji
#> 4         4  2.0   Average Fuji
#> 5         5  2.5   Average Fuji
#> 6         6  3.0   Average Fuji
```

```
tail(profiles)
```

```
#>  profileID price freshness      type
```

---

<sup>2</sup>Note that including hard constraints in your designs can substantially reduce the statistical power of your design, so use them cautiously and avoid them if possible.

```
#> 25      25    2.5 Excellent Honeycrisp
#> 26      26    3.0 Excellent Honeycrisp
#> 27      27    3.5 Excellent Honeycrisp
#> 28      28    4.0 Excellent Honeycrisp
#> 29      29    4.5 Excellent Honeycrisp
#> 30      30    5.0 Excellent Honeycrisp
```

## Generating designs

### Randomized designs

Once a set of profiles is obtained, a randomized conjoint survey can then be generated using the `cbc_design()` function:

```
design <- cbc_design(
  profiles = profiles,
  n_resp   = 900, # Number of respondents
  n_alts   = 3,   # Number of alternatives per question
  n_q      = 6    # Number of questions per respondent
)
```

```
dim(design) # View dimensions
```

```
#> [1] 16200      8
```

```
head(design) # Preview first 6 rows
```

```
#>  respID qID altID obsID profileID price      type freshness
#> 1      1  1  1      1      34    3.5      Gala    Average
#> 2      1  1  2      1       5    3.0      Fuji     Poor
#> 3      1  1  3      1      24    2.0      Fuji    Average
#> 4      1  2  1      2      43    1.0      Fuji  Excellent
#> 5      1  2  2      2      19    3.0 Honeycrisp     Poor
#> 6      1  2  3      2      28    4.0      Fuji    Average
```

For now, the `cbc_design()` function only generates a randomized design. Other packages, such as the `idefix` package, are able to generate other types of designs, including Bayesian D-efficient designs, and future versions of the `cbcTools` package will add a wrapper around the `idefix` package to integrate some of these features. The randomized design simply samples from the set of `profiles` while ensuring that no two profiles are the same in any choice question.

The structure of the resulting `design` data frame is such that each row specifies the levels shown for one alternative in a choice question. In addition to the attribute levels, the `design` data frame also includes the following variables for identifying different aspects of the survey:

- `respID`: Identifies each survey respondent.
- `qID`: Identifies the choice question answered by the respondent.

- **altID**: Identifies the alternative in any one choice observation.
- **obsID**: Identifies each unique choice observation across all respondents.
- **profileID**: Identifies the profile in **profiles**.

### Labeled designs (a.k.a. “alternative-specific” designs)

A “labeled” design (also known as an “alternative-specific” design) can also be generated. In labeled designs, the levels of one attribute are used as the labels for each alternative. To do so, the **label** argument can be set to one of the attributes. This by definition also sets the number of alternatives in each question to the number of levels in the chosen attribute, so the **n\_alts** argument is overridden. In the example below, the **type** attribute is used as the label:

```
design_labeled <- cbc_design(
  profiles = profiles,
  n_resp   = 900, # Number of respondents
  n_alts   = 3,   # Number of alternatives per question
  n_q      = 6,   # Number of questions per respondent
  label    = "type" # Set the "type" attribute as the label
)

dim(design_labeled)
```

```
#> [1] 16200      8
```

```
head(design_labeled)
```

```
#>  respID qID altID obsID profileID price      type freshness
#> 1      1  1  1      1         27  3.5      Fuji    Average
#> 2      1  1  2      1         55  3.5      Gala    Excellent
#> 3      1  1  3      1         39  2.5 Honeycrisp Average
#> 4      1  2  1      2         23  1.5      Fuji    Average
#> 5      1  2  2      2          9  1.5      Gala     Poor
#> 6      1  2  3      2         60  2.5 Honeycrisp Excellent
```

In the above example, the **type** attribute is now fixed to be the same order for every choice question, ensuring that each level in the **type** attribute will always be shown in each choice question.

### Adding a “no choice” option (a.k.a. “outside good”)

Often times designers may wish to allow respondents to opt out from choosing any of the alternatives shown in any one choice question. Such a “no choice” (or “outside good”) option can be included by setting **no\_choice = TRUE**. If included, all categorical attributes will be dummy-coded to appropriately dummy-code the “no choice” alternative.

```
design_nochoice <- cbc_design(
  profiles = profiles,
```



```

  n_resp    = 900, # Number of respondents
  n_alts    = 3, # Number of alternatives per question
  n_q       = 6, # Number of questions per respondent
  no_choice = TRUE
)

```

```
dim(design_nochoice)
```

```
#> [1] 21600    13
```

```
head(design_nochoice)
```

```

#>  respID qID altID obsID profileID price type_Fuji type_Gala type_Honeycrisp
#> 1      1  1  1      1         23   1.5          1          0              0
#> 2      1  1  2      1         47   3.0          1          0              0
#> 3      1  1  3      1         63   4.0          0          0              1
#> 4      1  1  4      1          0   0.0          0          0              0
#> 5      1  2  1      2         58   1.5          0          0              1
#> 6      1  2  2      2         20   3.5          0          0              1
#>  freshness_Poor freshness_Average freshness_Excellent no_choice
#> 1              0              1              0              0
#> 2              0              0              1              0
#> 3              0              0              1              0
#> 4              0              0              0              1
#> 5              0              0              1              0
#> 6              1              0              0              0

```

## Inspecting designs

The package includes some functions to quickly inspect some basic metrics of a design. The `cbc_balance()` function prints out a summary of the counts of each level for each attribute across all choice questions as well as the two-way counts across all pairs of attributes for a given design:

```
cbc_balance(design)
```

```

#> =====
#> price x type
#>
#>      Fuji Gala Honeycrisp
#>      NA 5496 5271      5433
#> 1   2315  783  786      746
#> 1.5 2305  788  744      773
#> 2   2302  797  746      759
#> 2.5 2265  767  705      793
#> 3   2364  804  768      792

```

```

#> 3.5 2347 790 793 764
#> 4 2302 767 729 806
#>
#> price x freshness
#>
#>      Poor Average Excellent
#>      NA 5373 5404 5423
#> 1 2315 779 778 758
#> 1.5 2305 770 786 749
#> 2 2302 746 784 772
#> 2.5 2265 775 729 761
#> 3 2364 782 768 814
#> 3.5 2347 769 781 797
#> 4 2302 752 778 772
#>
#> type x freshness
#>
#>      Poor Average Excellent
#>      NA 5373 5404 5423
#> Fuji 5496 1736 1911 1849
#> Gala 5271 1753 1726 1792
#> Honeycrisp 5433 1884 1767 1782

```

Similarly, the `cbc_overlap()` function prints out a summary of the amount of “overlap” across attributes within the choice questions. For example, for each attribute, the count under “1” is the number of choice questions in which the same level was shown across all alternatives for that attribute (because there was only one level shown). Likewise, the count under “2” is the number of choice questions in which only two unique levels of that attribute were shown, and so on:

```

cbc_overlap(design)

#> =====
#> Counts of attribute overlap:
#> (# of questions with N unique levels)
#>
#> price:
#>
#>   1    2    3
#> 82 1853 3465
#>
#> type:
#>
#>   1    2    3
#> 554 3622 1224
#>

```

```
#> freshness:
#>
#>      1      2      3
#> 531 3614 1255
```

## Simulating choices

Choices for a given `design` can be simulated using the `cbc_choices()` function. By default, random choices are simulated:

```
data <- cbc_choices(
  design = design,
  obsID = "obsID"
)

head(data)
```

#>	respID	qID	altID	obsID	profileID	price	type	freshness	choice
#> 1	1	1	1	1	34	3.5	Gala	Average	1
#> 2	1	1	2	1	5	3.0	Fuji	Poor	0
#> 3	1	1	3	1	24	2.0	Fuji	Average	0
#> 4	1	2	1	2	43	1.0	Fuji	Excellent	0
#> 5	1	2	2	2	19	3.0	Honeycrisp	Poor	0
#> 6	1	2	3	2	28	4.0	Fuji	Average	1

Choices can also be simulated according to an assumed prior model. The default model used is a multinomial logit model with fixed parameters. In the example below, the choices are simulated using a utility model with the following parameters:

- 1 continuous parameter for `price`
- 2 categorical parameters for `type` ("Gala" and "Honeycrisp")
- 2 categorical parameters for `freshness` ("Average" and "Excellent")

Note that for categorical variables (`type` and `freshness` in this example), the first level defined when using `cbc_profiles()` is set as the reference level. The example below defines the following utility model for simulating choices for each alternative  $j$ :

$$u_j = 0.1price_j + 0.1type_j^{\text{Gala}} + 0.2type_j^{\text{Honeycrisp}} + 0.1freshness_j^{\text{Average}} + 0.2freshness_j^{\text{Excellent}} + \varepsilon_j$$

```
data <- cbc_choices(
  design = design,
  obsID = "obsID",
  priors = list(
    price = 0.1,
    type = c(0.1, 0.2),
    freshness = c(0.1, 0.2)
  )
)
```

```
)
)
```

The prior model used for simulating choices can also include other, more complex models, such as models that include interaction terms or mixed logit models. For example, the example below is the same as the previous example but with an added interaction between **price** and **type**:

```
data <- cbc_choices(
  design = design,
  obsID = "obsID",
  priors = list(
    price = 0.1,
    type = c(0.1, 0.2),
    freshness = c(0.1, 0.2),
    `price*type` = c(0.1, 0.5)
  )
)
```

To simulate choices according to a mixed logit model where parameters follow a normal or log-normal distribution across the population, the **randN()** and **randLN()** functions can be used inside the **priors** list. The example below models the **type** attribute with two random normal parameters using a vector of means (**mean**) and standard deviations (**sd**) for each level of **type**:

```
data <- cbc_choices(
  design = design,
  obsID = "obsID",
  priors = list(
    price = 0.1,
    type = randN(mean = c(0.1, 0.2), sd = c(1, 2)),
    freshness = c(0.1, 0.2)
  )
)
```

## Analyzing power

The simulated choice data can be used to conduct a power analysis by estimating the same model multiple times with incrementally increasing sample sizes. As the sample size increases, the estimated coefficient standard errors will decrease (i.e. coefficient estimates become more precise), allowing the designer to identify the sample size required to achieve a desired level of precision.

The **cbc\_power()** function achieves this by partitioning the choice data into multiple sizes (defined by the **nbreaks** argument) and then estimating a user-defined choice model on each data subset. In the example below, 10 different sample sizes are used. All models are

estimated using the `logitr` package (Helveston 2021), and any arguments for estimating models with the `logitr` package can be passed through the `cbc_power()` function (see the `logitr` documentation for more details at <https://jhelvy.github.io/logitr/>)

```
power <- cbc_power(  
  data      = data,  
  pars      = c("price", "type", "freshness"),  
  outcome   = "choice",  
  obsID     = "obsID",  
  nbreaks   = 10,  
  n_q       = 6  
)
```

```
head(power)
```

```
#>   sampleSize      coef      est      se  
#> 1         90      price 0.01560102 0.05085851  
#> 2         90    typeGala -0.08013790 0.13171618  
#> 3         90  typeHoneycrisp 0.07418976 0.12945714  
#> 4         90  freshnessAverage -0.03428226 0.12957018  
#> 5         90  freshnessExcellent 0.04001875 0.12632890  
#> 6        180      price 0.00574527 0.03606545
```

```
tail(power)
```

```
#>   sampleSize      coef      est      se  
#> 45         810  freshnessExcellent -0.026874505 0.04278034  
#> 46         900      price -0.003997445 0.01641768  
#> 47         900    typeGala -0.064741539 0.04089035  
#> 48         900  typeHoneycrisp -0.007240872 0.04017063  
#> 49         900  freshnessAverage 0.024506658 0.04053370  
#> 50         900  freshnessExcellent -0.026598383 0.04057202
```

The `power` data frame contains the coefficient estimates and standard errors for each sample size. In the example above, it is clear that the standard errors for a sample size of 900 are much lower than those for a sample size of just 90.

Visualizing the results of the power analysis can be particularly helpful for identifying sample size requirements. Since the `cbc_power()` function returns a data frame in a “tidy” (or “long”) format (Wickham 2014), the results can be conveniently plotted using the popular `ggplot2` package (Wickham, Chang, and Wickham 2016). A `plot()` method is already included in `cbcTools` to create a simple `ggplot` of the power curves:

```
plot(power)
```

Of course, designers may be interested in aspects other than standard errors. By setting `return_models = TRUE`, the `cbc_power()` function will return a list of estimated models (one for each sample size increment), which can then be used to examine other model objects.

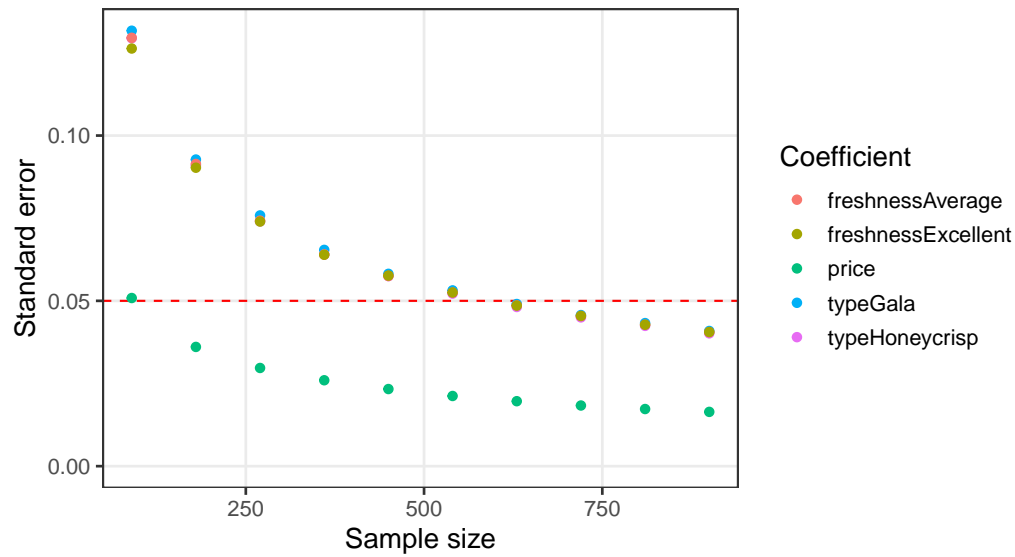


Figure 3: Standard errors of each model coefficient with increasing sample size.

The example below prints a summary of the last model in the list of returned models.

```
library(logitr)

models <- cbc_power(
  data      = data,
  pars      = c("price", "type", "freshness"),
  outcome   = "choice",
  obsID     = "obsID",
  nbreaks   = 10,
  n_q       = 6,
  return_models = TRUE
)

summary(models[[10]])

#> =====
#>
#> Model estimated on: Sat Jun 04 07:07:55 2022
#>
#> Using logitr version: 0.6.0
#>
#> Call:
#> FUN(data = X[[i]], outcome = ..1, obsID = ..2, pars = ..3, randPars = ..4,
#>      panelID = ..5, clusterID = ..6, robust = ..7, predict = ..8)
#>
#> Frequencies of alternatives:
#>      1      2      3
```

```

#> 0.33093 0.33389 0.33519
#>
#> Exit Status: 3, Optimization stopped because ftol_rel or ftol_abs was reached.
#>
#> Model Type:      Multinomial Logit
#> Model Space:      Preference
#> Model Run:        1 of 1
#> Iterations:        9
#> Elapsed Time:      0h:0m:0.03s
#> Algorithm:        NLOPT_LD_LBFGS
#> Weights Used?:     FALSE
#> Robust?            FALSE
#>
#> Model Coefficients:
#>
#>               Estimate Std. Error z-value Pr(>|z|)
#> price          -0.0039974  0.0164177 -0.2435  0.8076
#> typeGala        -0.0647415  0.0408904 -1.5833  0.1134
#> typeHoneycrisp  -0.0072409  0.0401706 -0.1803  0.8570
#> freshnessAverage  0.0245067  0.0405337  0.6046  0.5454
#> freshnessExcellent -0.0265984  0.0405720 -0.6556  0.5121
#>
#> Log-Likelihood:    -5.930144e+03
#> Null Log-Likelihood: -5.932506e+03
#> AIC:                1.187029e+04
#> BIC:                1.190326e+04
#> McFadden R2:        3.982252e-04
#> Adj McFadden R2:    -4.445889e-04
#> Number of Observations: 5.400000e+03

```

## Piping it all together!

One of the convenient features of how the package is written is that the object generated in each step is used as the first argument to the function for the next step. Thus, just like in the overall program diagram (see Figure 2), the functions can be piped together using either the Base R `|>` operator or the popular `%>%` operator from the `magrittr` package (Bache et al. 2022). The example below will generate the same power analysis plot as in Figure 3 by sequentially evaluating each of the main design steps.

```

cbc_profiles(
  price      = seq(1, 4, 0.5), # $ per pound
  type       = c('Fuji', 'Gala', 'Honeycrisp'),
  freshness  = c('Poor', 'Average', 'Excellent')
) |>
cbc_design(
  n_resp     = 900, # Number of respondents

```

```

  n_alts = 3,    # Number of alternatives per question
  n_q     = 6     # Number of questions per respondent
) |>
cbc_choices(
  obsID = "obsID",
  priors = list(
    price      = 0.1,
    type       = c(0.1, 0.2),
    freshness  = c(0.1, 0.2)
  )
) |>
cbc_power(
  pars      = c("price", "type", "freshness"),
  outcome   = "choice",
  obsID     = "obsID",
  nbreaks   = 10,
  n_q       = 6
) |>
plot()

```

One benefit of this piped structure is that it enables the designer to quickly observe the implications of different design choices on statistical power. For example, consider the following “what if” design questions:

- What if one more choice question was added to each respondent?
- What if the number of alternatives per choice question was decreased to two?
- What if a labeled design were used for the `type` attribute?
- What if there was an interaction effect between `price` and `type`?

Obtaining an answer to each of these questions requires only small modifications to the arguments in one or more functions inside the analysis “pipeline”. Once the change is made, the entire analysis can be quickly re-executed and the results compared to those of an alternative design. Because `cbcTools` leverages the `logitr` package (which is highly optimized for speed) for simulating choices and estimating models, re-executing each analysis should only seconds to compute the entire process on most modern computers and laptops.

## Conclusions

This paper introduces an open source R package, `cbcTools`, for generating and evaluating choice-based conjoint experiment designs. The package contains functions for generating experiment designs, examining attribute balance and overlap, simulating choice data, and conducting power analyses. In addition to being open source, one of its primary advantages over alternatives is the ability to examine the statistical power of different designs under a variety of conditions, such as when preference heterogeneity or strong interactions between certain attributes may be expected in respondent choices. Users can simulate choice data



for designs according to multinomial and mixed logit models and examine sample size requirements to achieve a desired level of precision for multinomial and mixed logit models. The package is designed such that each function along the analysis process generates an object for the next function, enabling the ability to conveniently pipe together a full analysis from defining attributes and levels, generating an experiment design, simulating choice data, and examining statistical power. Detailed package documentation can be found at <https://jhelvy.github.io/cbcTools/>.

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