# Validate Power: Westfall-Young

December 29, 2021

### Explanation

#### Westfall-Young procedures

The Westfall-Young procedure was validated separately due to its unique complications and computational burden.

First, we wrote a series of careful unit tests ensuring that our code worked as expected for each step of the WY procedure. One of these tests compared results from our procedure to the WY procedure in the multtest package, and found it matched quite well.

Second, we also tested the WY procedures using the same simulation procedure described above. For constant effect and fixed effect models, the power estimation matches between PUM and the simulations. However, for random effects models, the two methods diverge in some cases. We find that the simulated power matches the PUMP-calculated power only when (1) there is a large number of blocks/clusters, and (2) the user has a large number of WY permutations. Below, we discuss the single-step procedure because it is simpler, although the same concepts hold for the step-down procedure. Although it is difficult to verify why this discrepancy occurs, our hypothesis is that this behavior occurs due to the combination of the sensitivity of the WY procedure, and the instability of the random effects model.

The goal of the WY procedure is to estimate the adjusted p-value for outcome i

$$\tilde{p}_i = Pr\left(\min_{1 \le j \le k} P_j \le p_i \mid H_0^C\right).$$

where  $P_j$  is a random variable representing a p-value for outcome j, and lowercase  $p_i$  is the observed realization for outcome i. We assume a set of k tests with corresponding null hypotheses  $H_{0i}$  for  $i=1,\dots,k$ . The complete null hypothesis is the setting where all the null hypotheses are true:  $H_0^C = \bigcap_{i=1}^k H_{0i} = \{\text{all } H_i \text{ are true}\}$ . In order to estimate this p-value, the p-value is calculated across B permutations

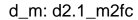
$$\tilde{p}_i = \frac{1}{B} \sum_{b=1}^{B} \mathbb{1} \left( \min_{1 \le j \le k} p_{b,j}^* \le p_i \right).$$

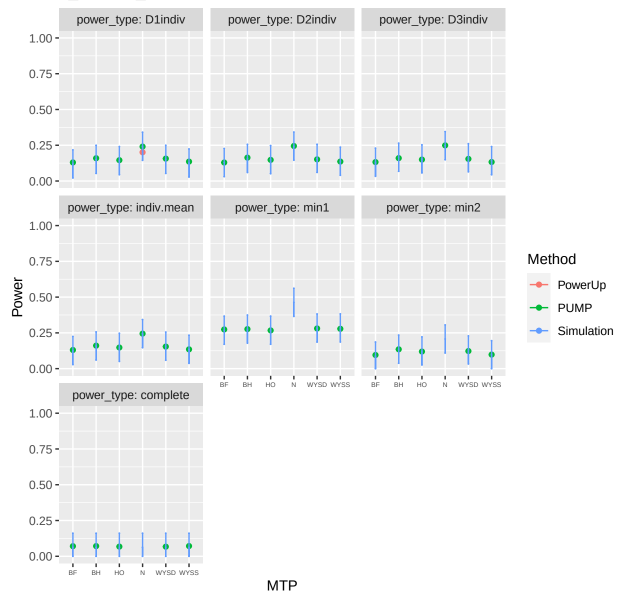
In order for the procedure to be correct, we should be generating the p-values  $P_j$  from the the true distribution of p-values under the null hypothesis. We generally are testing outcomes that are truly significant, so the observed p-values are small. Thus, looking at these expressions, we are concerned about tail behavior; we are testing whether a small observed p-value is less than the minimum of of the generated p-values of our multiple outcomes. This combination of factors means that a small discrepancy in the tails between the generated null distribution and the true distribution could have a large impact on the adjusted p-value.

Given that we are relying on tail behavior, this explains why we need both a large number of permutations, and a large number of blocks/clusters. The large number of permutations is required because we are trying to estimate a rare event. With a significant outcome, it is rare that the observed p-value will be less than the minimum of p-values generated from the null distribution. The large number of blocks/clusters is required to accurately estimate the tail behavior for random effects models. With a small number of blocks/clusters, we may not properly estimate the spread of the distribution, resulting in a poor estimate of the tail behavior.

## d2.1 models

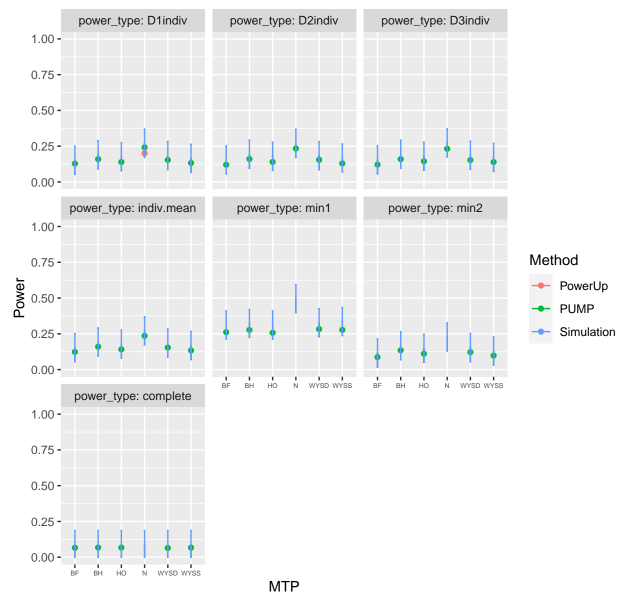
Constant effects





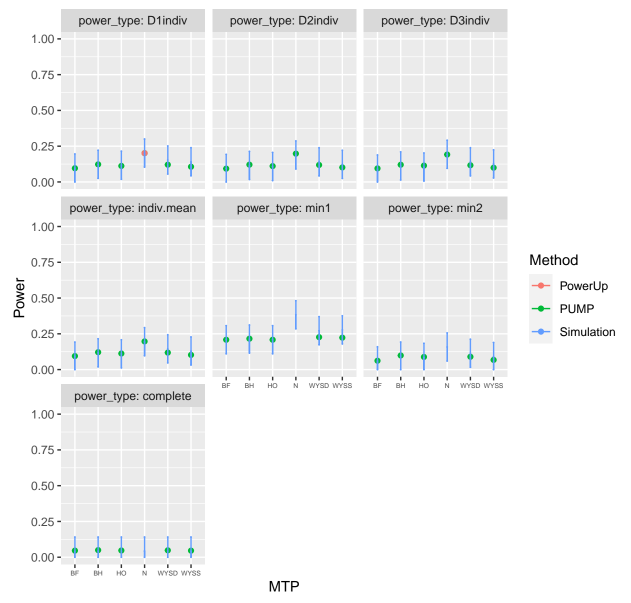
### Fixed effects

## d\_m: d2.1\_m2ff



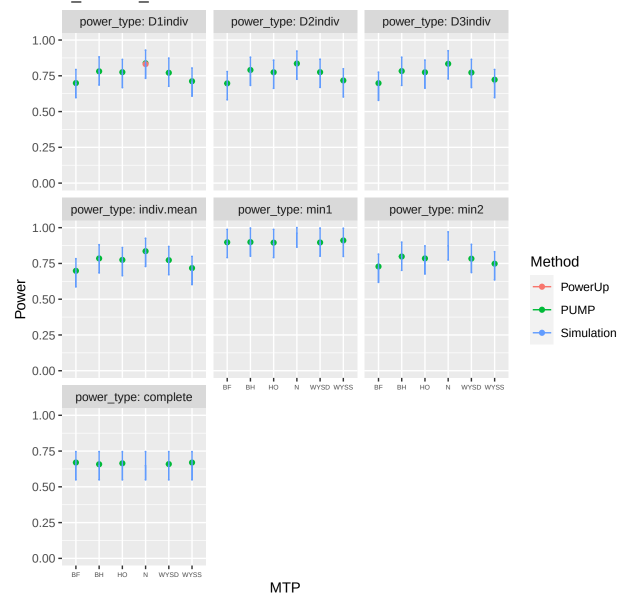
### Random effects

## d\_m: d2.1\_m2fr



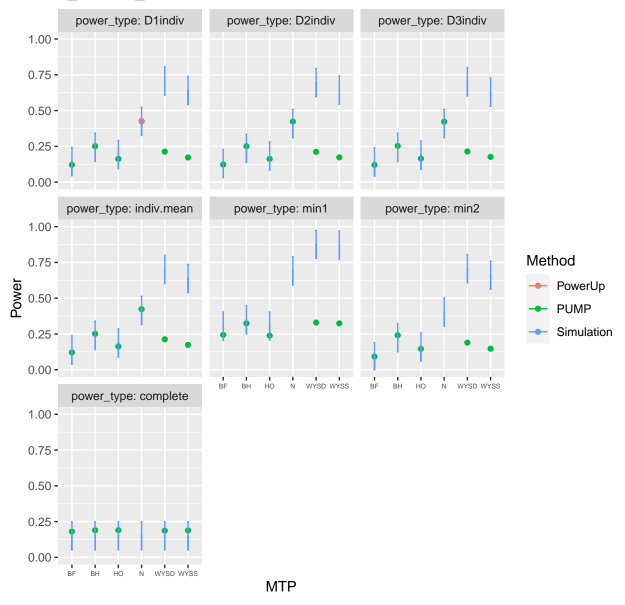
## d2.2 models

## d\_m: d2.2\_m2rc



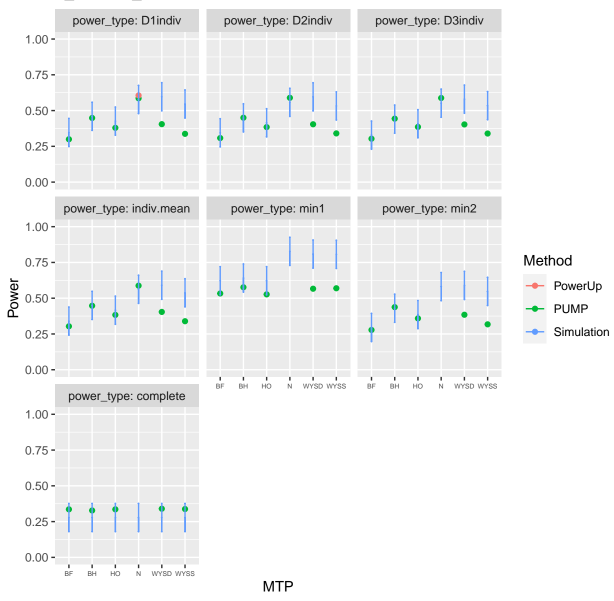
## d3.1 models

d\_m: d3.1\_m3rr2rr



## d3.3 models

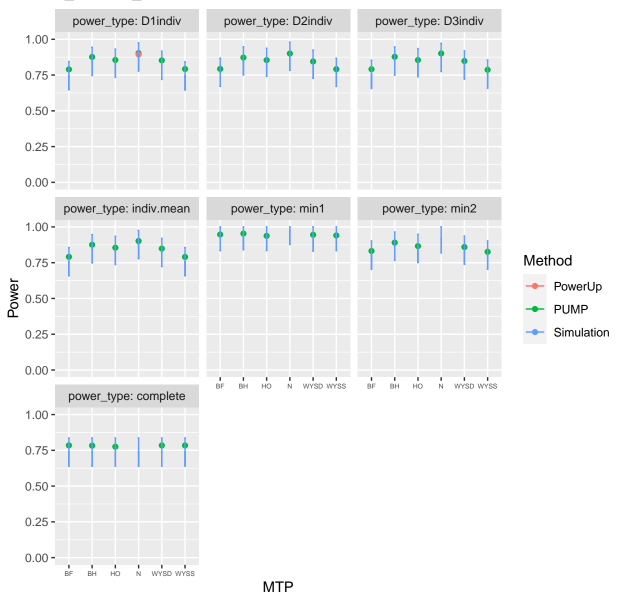
d\_m: d3.3\_m3rc2rc



## d3.2 models

Constant effects

## d\_m: d3.2\_m3ff2rc



## Random effects

## d\_m: d3.2\_m3rr2rc

