# Verification Report: Immunogenicity Graphs

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# Description

This is the verification report for the immuno\_graphical folder of the correlates\_reporting project for CoVPN.

In the verification process, the outputs of immuno\_graphical\_verification\_code.R were visually compared against the plots listed in the file immuno\_graphical/verification/verification\_plan.pdf. The values presented in the original plots were also generated independently by the tester and compared against the outputs generated by the original programmer.

The files immuno\_graphical/verification/verification\_input/practice\_data.csv, immuno\_graphical/verification and immuno\_graphical/verification/long\_twophase\_data.csv were provided by the original programmer to the tester for verification purposes.

### **Data Verification**

Load dat.twophase.sample

```
verification.dat.twophase.sample <- read.csv(
  here("verification/output/dat.twophase.sample_verification.csv")
)
original.dat.twophase.sample<-read.csv(
  here("verification/input/twophase_data.csv")
)</pre>
```

# Verification

```
data_clean_comparison <- compare_datasets(
  cols = colnames(verification.dat.twophase.sample), index = "Ptid",
  ds1 = verification.dat.twophase.sample, ds2 = original.dat.twophase.sample)</pre>
```

## There are 0 mismatched fields of 83.

dat.twophase.sample generated by using immuno\_graphical\_verification\_code.R was equivalent to the output generated by original programmer, and passes verification.

#### Load dat.long.twophase.sample

```
verification.dat.long.twophase.sample <- read.csv(
  here("verification/output/dat.long.twophase.sample_verification.csv")
)
original.dat.long.twophase.sample <- read.csv(
  here("verification/input/long_twophase_data.csv")
)</pre>
```

#### Verification

```
data_clean_comparison <- compare_datasets(
  cols = colnames(verification.dat.long.twophase.sample), index = "Ptid",
  ds1 = verification.dat.long.twophase.sample, ds2 = original.dat.long.twophase.sample)</pre>
```

## There are 0 mismatched fields of 39.

dat.long.twophase.sample generated by using immuno\_graphical\_verification\_code.R was equivalent to the output generated by original programmer, and passes verification.

## Verification of pair plots of D57 Ab markers: baseline negative vaccine arm

```
data_clean_comparison <- compare_datasets(
  cols = colnames(verification_spearman_correlation),index = "corr",
  ds1 = verification_spearman_correlation, ds2 = original_spearman_correlation
)</pre>
```

## There are 0 mismatched fields of 3.

The scatter plots and density plots generated by the teseter were visually compared with the outputs generated by original programmer, the partial Spearman's correlations calculated by tester were equivalent to the partial Spearman's correlations provided by original programmer, and passes verification.

# Verification of RCDF plots for D57 Ab markers: by baseline status for the vaccine arm.

```
data_clean_comparison <- compare_datasets(
  cols = colnames(verification_wrcdf), index = "x",
  ds1 = verification_wrcdf, ds2 = original_wrcdf
)</pre>
```

## There are 0 mismatched fields of 9.

RCDF plot generated by tester was visually compared with the RCDF plot generated by the original programmer. Key statistics(0th, 10th, 20th, 30th, 40th, 50th, 60th, 70th, 80th, 90th, 100th percentile) of RCDF calculated by tester were equivalent to the key statistics provided by original programmer, and passes verification.

Verification of boxplots of D57 Ab markers: baseline negative vaccine + placebo arms

```
data_clean_comparison <- compare_datasets(
  cols = colnames(original_boxplot), index = "summary",
  ds1 = original_boxplot, ds2 = verification_boxplot
)</pre>
```

## There are 0 mismatched fields of 9.

Boxplots generated by tester were visually compared with the boxplots generated by the original programmer. Key statistics(min, 25th percentile, median, mean, 75th percentile, max) showed in the boxplot calculated by tester were equivalent to the key statistics provided by original programmer, and passes verification.

Verification of spaghetti plots of Ab markers over time: baseline negative vaccine + placebo arm

```
data_clean_comparison <- compare_datasets(
  cols = colnames(verification_spaghetti_plot_data_BaselineNeg_BaselinePos), index = "Ptid",
  ds1 = verification_spaghetti_plot_data_BaselineNeg_BaselinePos, ds2 = original_spaghetti_plot_data_BaselineNeg_BaselinePos</pre>
```

## There are 0 mismatched fields of 7.

Spaghetti plots generated by tester were visually compared with the Spaghetti plots generated by the original programmer. Input data for the Spaghetti plots generated by tester was equivalent to the input data provided by original programmer, and passes verification.

# Signature

| Role   | Name  | Signature | Date |
|--------|-------|-----------|------|
| Tester | Di Lu |           |      |