Stokes Radius Analysis by Gel Filtration

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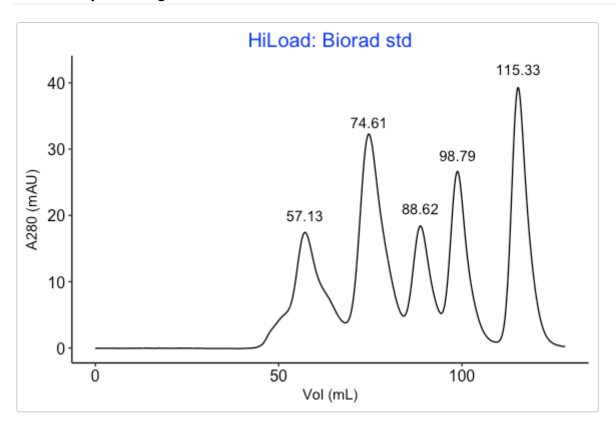
2016-09-02

```
## Loading required package: ggplot2

##
## Attaching package: 'ggplot2'

## The following object is masked from 'package:radii':
##
## ggplot
```

Load and plot the gel filtration data



Transform the standard and then fit to the Stokes radius model

```
# superdex 200 HiLoad 16/60
vt = mean(c(122.31,122.93))
vo = mean(c(47.26,46.59))
```

name	mw	radii	ve	sigma	erfcinv
vitamin B12	1350	NA	115.33	0.9036924	0.0855587
horse myoglobin	17000	1.89	98.79	0.6851840	0.2866607
chicken ovalbumin	44000	2.75	88.62	0.5508290	0.4218022
bovine gamma-globulin	158000	5.22	74.61	0.3657441	0.6395597
bovine thyroglobulin	670000	8.50	57.13	0.1348174	1.0573877

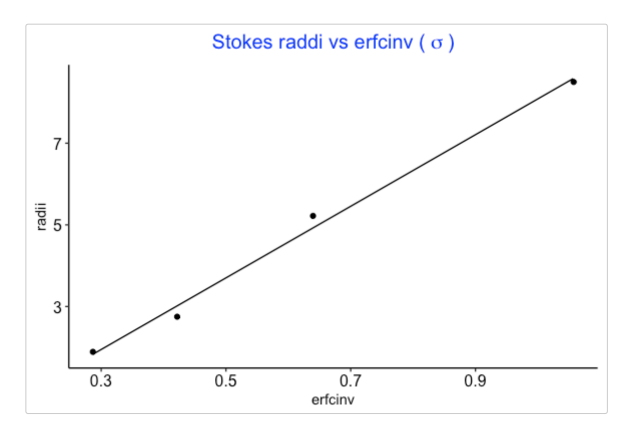
```
protein_std_HiLoad <- na.omit(protein_std_HiLoad)

fit <- lm (radii~erfcinv+1, data = protein_std_HiLoad)
summary(fit)</pre>
```

```
##
## Call:
## lm(formula = radii ~ erfcinv + 1, data = protein_std_HiLoad)
## Residuals:
##
         2
                  3
  0.05920 -0.26571 0.29500 -0.08848
##
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.6826 0.3329 -2.051 0.17676
                       0.4980 17.608 0.00321 **
## erfcinv
                8.7679
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.2907 on 2 degrees of freedom
## Multiple R-squared: 0.9936, Adjusted R-squared: 0.9904
## F-statistic: 310 on 1 and 2 DF, p-value: 0.00321
```

```
xdat <- seq(min(protein_std_HiLoad$erfcinv), max(protein_std_HiLoad$erfcinv),by = 0.01 )
pred <- predict(fit, newdata = data.frame(erfcinv=xdat))

# Plotting
g = radii:::ggplot() + geom_point(data = protein_std_HiLoad, aes(erfcinv, radii))
g <- g + ggtitle(expression("Stokes raddi vs erfcinv (" ~ sigma ~ ")"))
g + geom_line(aes(xdat,pred))</pre>
```



Purified VWF Propeptide

```
# superdex 200 HiLoad 16/60
vt = mean(c(122.31,122.93))
vo = mean(c(47.26,46.59))
# sample data
ve_D1D2_pure = 74.36
sigma_D1D2_pure=radii::partition_coef(ve_D1D2_pure, vo=vo, vt=vt)
erfcinv_D1D2_pure= radii::erfcinv(sigma_D1D2_pure)
r_D1D2_pure = predict(fit, newdata = data.frame(erfcinv=erfcinv_D1D2_pure))
print(paste("Stokes radius of the purified D1D2", r_D1D2_pure, "nm"))
```

```
## [1] "Stokes radius of the purified D1D2 4.9637398754402 nm"
```

Plasma D1D2

```
# D1D2
ve_D1D2_plasma = 81
sigma_D1D2_plasma=radii::partition_coef(ve_D1D2_plasma, vo=vo, vt=vt)
erfcinv_D1D2_plasma= radii::erfcinv(sigma_D1D2_plasma)
r_D1D2_plasma = predict(fit, newdata = data.frame(erfcinv=erfcinv_D1D2_plasma))
print(paste("Stokes radius of the purified D1D2", r_D1D2_plasma, "nm"))
```

[1] "Stokes radius of the purified D1D2 3.99917971664297 nm"

```
summary(fit)
```

```
##
## Call:
## lm(formula = radii ~ erfcinv + 1, data = protein_std_HiLoad)
##
## Residuals:
         2
                3 4
##
## 0.05920 -0.26571 0.29500 -0.08848
##
## Coefficients:
            Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -0.6826 0.3329 -2.051 0.17676
## erfcinv 8.7679 0.4980 17.608 0.00321 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.2907 on 2 degrees of freedom
## Multiple R-squared: 0.9936, Adjusted R-squared: 0.9904
## F-statistic: 310 on 1 and 2 DF, p-value: 0.00321
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